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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶:
A61K 31/41, 31/415, 31/44, 31/47, 31/505, 38/04

(11) International Publication Number:

WO 98/30216

A1 |

(43) International Publication Date:

16 July 1998 (16.07.98)

(21) International Application Number:

PCT/US98/00534

(22) International Filing Date:

7 January 1998 (07.01.98)

(74) Common Representative: MERCK & CO., INC.; 126 East Lincoln Avenue, Rahway, NJ 07065 (US).

(30) Priority Data:

60/034,927 9704197.4 10 January 1997 (10.01.97) U 28 February 1997 (28.02.97)

US GB

(71) Applicant (for all designated States except US): MERCK & CO., INC. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): BEERE, Polly, A. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). CHANG, Paul, I. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). PITT, Bertram [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). RUCINSKA, Ewa, J. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). SEGAL, Robert [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). SHARMA, Divakar [IN/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US). SNAVELY, Duane, B. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065 (US).

(81) Designated States: AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CU, CZ, EE, GE, GW, HU, ID, IL, IS, JP, KG, KR, KZ, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, SL, TJ, TM, TR, TT, UA, US, UZ, VN, YU, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: USE OF ANGIOTENSIN II ANTAGONISTS TO TREAT SYMPTOMATIC HEART FAILURE

(57) Abstract

Angiotensin II receptor antagonists are useful in reducing and preventing mortality and sudden cardiac death in symptomatic heart failure patients. Losartan potassium has been shown to reduce mortality and sudden cardiac death in this patient population. Additionally, losartan potassium has been shown to reduce the need for hospitalization of symptomatic heart failure patients.

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TITLE OF THE INVENTION

USE OF ANGIOTENSIN II ANTAGONISTS TO TREAT SYMPTOMATIC HEART FAILURE

5 BACKGROUND OF THE INVENTION

Angiotensin-converting-enzyme (ACE) inhibitors have been shown to reduce morbidity and mortality in patients with chronic heart failure and systolic left ventricular dysfunction as well as in patients post myocardial infarction. (See The CONSENSUS Trial Study Group.

- 10 Effects of enalapril on mortality in severe congestive heart failure.

 Results of the Cooperative North Scandinavian Enalapril Survival Study (CONSENSUS). N Engl J Med 1987; 316: 1429-1435; The SOLVD Investigators. Effect of enalapril on survival in patients with reduced left ventricular ejection fractions and congestive heart failure. N Engl J
- 15 Med 1991; 325: 293-302; The SOLVD Investigators. Effect of enalapril on mortality and the development of heart failure in asymptomatic patients with reduced left ventricular ejection fractions. N Engl J Med 1992; 327: 685-691; Cohn JN, Johnson G, Ziesche S, et al. A comparison of enalapril with hydralazine-isosorbide dinitrate in the
- treatment of chronic congestive heart failure. N Engl J Med 1991; 325: 303-310; Pfeffer MA, Braunwald E, Moye LA, et al. on behalf of the SAVE Investigators. Effect of captopril on mortality and morbidity in patients with left ventricular dysfunction after myocardial infarction. Results of the Survival and Ventricular Enlargement Trial. N Engl J
- 25 Med 1992; 327: 669-677; The Acute Infarction Ramipril Efficacy (AIRE) Study Investigators. Effect of ramipril on mortality and morbidity of survivors of acute myocardial infarction with clinical evidence of heart failure. Lancet 1993; 342: 812-828; Fonarow GC, Chelimsky-Fallick C, Warner Stevenson L, et al. Effect of direct
- vasodilation with hydralazine versus angiotensin-converting enzyme inhibition with captopril on mortality in advanced heart failure: the Hy-C trial. J Am Coll Cardiol 1992; 19: 842-850; Gruppo Italiano per lo Studio delia Sopravvivenza nell'infarto Miocardico. GISSI-3: Effects of lisinopril and transdermal glyceryl trinitrate singly and together on 6-
- 35 week mortality and ventricular function after acute myocardial

infarction. Lancet 1994; 343: 1115-1122; ISIS Collaborative Group OU. ISIS-4: Randomized study of oral isosorbide mononitrate in over 50,000 patients with suspected acute myocardial infarction. Circulation 1993; 88: I394.) The benefits of ACE inhibitors have been attributed to blockade of angiotensin II production and/or to a decrease in the breakdown of bradykinin. (See Pitt B, Chang P, Timmermans P. Angiotensin II receptor antagonists in heart failure: Rationale and design of the Evaluation of Losartan in the Elderly (ELITE) Trial. Cardiovascular Drugs and Therapy 1995; 9: 693-700; and Gavras I. Bradykinin-mediated effects of ACE inhibition. Kidney Int 1992; 42: 10 1020-1029.) Bradykinin has been shown to have beneficial effects associated with the release of nitric oxide and prostacyclin which may contribute to the hemodynamic effects of ACE inhibition. Bradykinin may, however, also be responsible for certain adverse effects associated with use of ACE inhibitors, such as cough, angioedema, renal 15 dysfunction, and hypotension. (See Pitt B, Chang P, Timmermans P. Angiotensin II receptor antagonists in heart failure: Rationale and design of the Evaluation of Losartan in the Elderly (ELITE) Trial. Cardiovascular Drugs and Therapy 1995; 9: 693-700; Gavras I. Bradykinin-mediated effects of ACE inhibition. Kidney Int 1992; 42: 20 1020-1029: Israili ZH, Hall WD. Cough and angioneurotic edema associated with angiotensin-converting enzyme inhibitor therapy. A review of the literature and pathophysiology. Ann Intern Med 1992; 117: 234-242; Chalmers D, Dombey SL, Lawson DH. Post-marketing surveillance of captopril (for hypertension): a preliminary report. Br J 25 Clin Pharmacol 1987; 24: 343-349; Lacourciere Y, Brunner H, Irwin R, et al. and the Losartan Cough Study Group. Effects of modulators of the renin-angiotensin-aldosterone system on cough. J Hypertension 1994; 12: 1387-1393.) These adverse effects may account in part for the fact that ACE inhibitors are used in less than 30 percent of patients 30 with heart failure in spite of their proven clinical benefit. (See Stafford RS, Saglam D, Blumenthal D. Low rates of angiotensin-converting enzyme inhibitor use in congestive heart failure. Circulation 1996; 94: I-194(Abstract)).

The development of orally-active, nonpeptidic angiotensin II type 1 receptor antagonists such as losartan, has provided the opportunity to block the angiotensin II type 1 receptor specifically without increasing bradykinin levels. (See Timmermans P, Wong PC, Chiu AT, et al. Angiotensin II receptors and angiotensin II receptor antagonists. Pharmacol Reviews 1993; 45: 205-251.) Since angiotensin

- antagonists. Pharmacol Reviews 1993; 45: 205-251.) Since angiotensin II may be produced by alternate pathways, losartan may offer additional advantages over treatment with ACE inhibitors where blockade of the effects of angiotensin II may be incomplete. (See Miura S, Ideishi M,
- 10 Sakai T, et al. Angiotensin II formation by an alternative pathway during exercise in humans. J Hypertension 1994; 12: 1177-1181; Urata H, Kinoshita A, Misono KS, Bumpus FM, Husain A. Identification of a highly specific chymase as the major angiotensin II-forming enzyme in the human chymase. J Biol Chem 1990; 265: 22348-22357; Urata H,
- 15 Strobel F, Ganten D. Widespread tissue distribution of human chymase. J Hypertension 1994; 12: S17-S22; Aldigier JC, Huang H, Dalmay F, et al. Angiotensin-converting enzyme inhibition does not suppress plasma angiotensin II increase during exercise in humans. J Cardiovasc Pharmacol 1993; 21: 289-295.) Losartan is indicated for the treatment
- of hypertension in many countries and in earlier studies in patients with symptomatic heart failure, oral losartan produced beneficial hemodynamic effects both acutely and with chronic dosing. (See Crozier I, Ikram H, Awan N, et al. Losartan in heart failure: Hemodynamic effects and tolerability. Circulation 1995; 91: 691-697;
- and Gottlieb SS, Dickstein K, Fleck E, et al. Hemodynamic and neurohormonal effects of the angiotensin II antagonist losartan in patients with congestive heart failure. Circulation 1993; 88: 1602-1609.)

The Evaluation of Losartan In The Elderly (ELITE) Study was conducted to compare effects on renal function, morbidity/ mortality, and tolerability of long-term treatment with losartan versus captopril, in older patients with symptomatic heart failure.

SUMMARY OF THE INVENTION

30

A method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of

an angiotensin II antagonist. A method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist. A method for reducing mortality and sudden cardiac death by

5 administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist is disclosed. A method for preventing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist. A method for reducing hospitalization by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1. ELITE Study Profile.

- 15 (* While on study therapy; and ** Withdrawn from assigned therapy, but still followed up for the intention-to-treat analysis of secondary endpoint.)
- Figure 2. Kaplan-Meier Survival Curves among Patients with Chronic
 Heart Failure in the Losartan and Captopril Groups.

 (Patients in the losartan group had a 46 percent lower risk of death
 (RR= risk reduction) than patients in the captopril group (P=0.035).

 Patients were followed for 48 weeks.)
- Figure 3. Effect of Losartan on Death in Various Subgroups.

 (**One losartan-treated patient did not have EF measured; and **E**

 Based on patient history. For each subgroup, the percentage of risk reduction (RR) with losartan is plotted (solid squares). Horizontal lines represent 95 percent confidence intervals. The size of each square is proportional to the percentage of events in the subgroup. The light square at the bottom of the panel represents the overall result for death. The bold vertical line corresponds to a finding of no effect. The RRs in individual subgroups are generally consistent with the overall RR, except for females. Interaction tests for the effect of losartan on mortality in gender was significant at the 10 percent level (p=0.053).

Tests for interaction in the other subgroups were not significant. NYHA denotes New York Heart Association.)

Figure 4. Changes in New York Heart Association Functional Class.

(*P≤0.001. Percent of patients with change in New York Heart Association (NYHA) Functional Class at week 48 versus Baseline.)

Figure 5. Comparison of twelve-week mortality data from the SOLVD study, the losartan exercise studies (U.S. and international combined), and the ELITE study.

(pbo = placebo; enal = enalapril; los = losartan; capt = captopril).

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DETAILED DESCRIPTION OF THE INVENTION

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This invention concerns a method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist.

An embodiment of this invention is the method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720,

- 25 LRB081, SC52458, SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492,
- 30 YH1498, and YM31472. The preferred angiotensin II receptor antagonists useful in this method are: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

Another embodiment of the invention is the method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of an imidazole angiotensin II receptor antagonist of formula I

5

wherein:

R¹ is:

R2 is H; Cl; Br; I; F; NO2; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO2H; CO2R9; HNSO2CH3; NHSO2CF3; CONHOR¹²; SO_2NH_2 ; N-N; aryl; or furyl; N

5

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

10

- R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;
- R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or 15 CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl or cycloalkylalkynyl 5 to 10 carbon atoms; (CH2)_sZ(CH2)_mR⁵ optionally substituted with F or CO₂R₁₄; benzyl substituted on the phenyl ring with 1 or 2 20 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

 R^7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where $\nu=1-6$, C₆F₅; CN;

25

—Ü−R¹⁶ ; straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

	R ⁸ is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH ₂) _m -imidazol-1-yl; -(CH ₂) _m -1,2,3-triazolyl
5	optionally substituted with one or two group selected from CO ₂ CH ₃ or alkyl of 1 to 4 carbon atoms; -(CH ₂) _s
	tetrazolyl;
10	O -(CH ₂) _{n-1} CH-R ¹¹ ; -(CH ₂) _n OCR ¹⁴ ; -(CH ₂) _n SR ¹⁵ ; OR ¹⁷
	R ¹⁴ O O -CH=CH(CH ₂) ₅ CHOR ¹⁵ ; -CH=CH(CH ₂) ₅ CR ¹⁶ ; -CR ¹⁶ ;
15	-CH=CH(CH ₂) ₅ OCR ¹¹ ; (CH ₂) ₅ -CH-COR ¹⁶ ; CH ₃
	O Y Y -(CH ₂) _n CR ¹⁶ ; -(CH ₂) _n OCNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ COR ¹⁰ ;
•	
	Q -(CH ₂) _n NR ¹¹ CNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ SO ₂ R ¹⁰ ;
20	Y -(CH ₂) _n NR ₁₁ CR ₁₀ ; -(CH ₂) _m F; -(CH ₂) _m ONO ₂ ; -CH ₂ N ₃ ; -(CH ₂) _m NO ₂ ; -CH=N-NR ₁₁ R ₁₇ ;

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N$$

- 5 R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)_pC6H5;
- R¹¹ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

R¹⁴ is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

5

- R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
- 10 R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;
 - R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

15

R¹⁸ and R¹⁹ independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 Q ;

- Q is NR²⁰, O or CH₂;
- R20 is H, alkyl of 1-4 carbon atoms, or phenyl;
- 25 R²¹ is alkyl of 1 to 6 carbon atoms, -NR²²R²³, or CHCH₂CO₂CH₃; NH₂

R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

5 R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

—
$$NHSO_2$$
—CH₃; or — $NHSO_2$ —

- 10 R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
 - R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;
- 15 R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)_q-;
 - R³¹ is H, alkyl or 1 to 4 carbon atoms, -CH₂CH=CH₂ or -CH₂C₆H₄R³²;

20

X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-,

R26 R23 R23

-NHC(R²⁷)(R²⁸)-, -NR²³SO₂-, -SO₂NR²³-, -CH=CH-, -

CF=CF-, -CH=CF-, -CF=CH-, -CH2CH2-, -C(R27)(R28)NH-,

Y is O or S;

Z is O, NR11, or S;

m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

5 q is 2 to 3;

r is 0 to 2;

s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds;

10 provided that:

- (1) the R¹ group is not in the ortho position;
- (2) when R¹ is

$$-x - \begin{cases} R^{13} \\ -x - R^{3} \end{cases}$$

15

X is a single bond, and R13 is CO₂H, or

20

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

(3) when R1 is

$$-x - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

5
(4) when R¹ is 4-CO₂H or a salt thereof, R⁶ cannot be S-alkyl;

(5) when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;

(6) when R¹ is

10

20

15 X is -OCH₂-, and R^{13} is 2-CO₂H, and R^{7} is H then R^{6} is not C₂H₅S;

(7) when R^1 is

- 14 -

and R6 is n-hexyl then R7 and R8 are not both hydrogen;

(8) when R1 is

5

R6 is not methoxybenzyl;

10 (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;

(10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

X is -NH-C-, R¹³ is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^{3} \end{array} \right)$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

10

$$R^1 = X - \begin{pmatrix} & & & \\$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right),$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 4-(tetrazol-5-yl).

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A preferred embodiment of the invention is the method as recited above, using the imidazole angiotensin II receptor antagonist of formula I:

5

wherein:

R1 is -CO2H; -NHSO2CF3;

10

R6 is alkyl of 3 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, alkynyl of 3 to 10 carbon atoms, cycloalkyl of 3 to 8 carbon atoms, benzyl substituted on the phenyl ring with up to two groups selected from alkoxy of 1 to 4 carbon atoms, halogen, alkyl of 1 to 4 carbon atoms, and nitro;

15

R8 is phenylalkenyl wherein the aliphatic portion is 2 to 4 carbon atoms, $-(CH_2)_m$ -imidazol-1yl, $-(CH_2)_m$ 1,2,3-triazolyl optionally substituted with one or two groups selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms,

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(CH_2)_m-tetrazolyl, -(CH_2)_nOR^{11}; -(CH_2)_nO^{\bar{1}}_{CR}14;
                       Q R14
-CH=CH(CH2)sCR16, -CH=CH(CH2)sCHOR15;
 5
                       O O -(CH<sub>2</sub>)<sub>n</sub>CR<sup>16</sup>; -(CH<sub>2</sub>)<sub>n</sub>NHCOR<sup>10</sup>; -(CH<sub>2</sub>)<sub>n</sub>NHSO<sub>2</sub>R<sup>10</sup>;
                       _{-(CH_2)_mF;}^{O}
               R13 is -CO<sub>2</sub>H, -CO<sub>2</sub>R<sup>9</sup>, NHSO<sub>2</sub>CF<sub>3</sub>; SO<sub>3</sub>H;
10
                       or N-N
               R16 is H, alkyl of 1 to 5 carbon atoms, OR17, or NR18R19;
               X is carbon-carbon single bond, -CO-, -CON-, -CH2CH2-, -NCO-, R23 R23
15
                       -OCH2-, -CH2O-, -SCH2-, -CH2S-, -NHCH2-, -CH2NH- or
                       -CH=CH-; and pharmaceutically acceptable salts of these
                       compounds.
20
                       A further preferred embodiment of the invention is the
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A further preferred embodiment of the invention is the method as recited above, using the imidazole angiotensin II receptor antagonist of formula I:

R² is H, alkyl of 1 to 4 carbon atoms, halogen, or alkoxy of 1 to 4 carbon atoms;

R6 is alkyl, alkenyl or alkynyl of 3 to 7 carbon atoms;

R7 is H, Cl, Br,
$$C_{\nu}F_{2\nu+1}$$
, where v=1-3, or -CR16;

30

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O R¹⁴ R⁸ is -(CH₂)
$$_m$$
OCR¹¹; -(CH₂) $_m$ OCR¹⁴; -CH=CH-CHOR¹⁵;

$$\begin{array}{c} O & O \\ -(CH_2)_m CR^{16}; -CH_2NHCOR^{10}; \\ -(CH_2)_m NHSO_2 R^{10}; & N-N \\ -CH_2 & N \end{array}; \text{ or } -COR^{16}; \end{array}$$

5 R10 is CF3, alkyl of 1 to 6 carbon atoms or phenyl;

R11 is H, or alkyl of 1 to 4 carbon atoms;

R13 is CO₂H; CO₂CH₂OCOC(CH₃)₃; NHSO₂CF₃;

and
$$N-N$$

10

15

R¹⁴ is H, or alkyl of 1 to 4 carbon atoms;

R¹⁵ is H, alkyl of 1 to 4 carbon atoms, or acyl of 1 to 4 carbon atoms;

R16 is H, alkyl of 1 to 5 carbon atoms; OR17; or

$$-N$$
 $O;$

20

m is 1 to 5;

X is single bond, -O-; -CO-; -NHCO-; or -OCH2-; and pharmaceutically acceptable salts.

A preferred embodiment of this invention is the method as recited above, using an imidazole angiotensin II receptor antagonist selected from the group consisting of:

2-Butyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.

- 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 5 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-[(methoxy-carbonyl)aminomethyl]imidazole.
 - 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-[(propoxy-carbonyl)aminomethyl]imidazole.
 - 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]imidazole-5-carboxaldehyde.
 - 2-Butyl-1-[(2'-carboxybiphenyl-4-yl)methyl]-imidazole-5-carbox-aldehyde.
 - 2-(1E-Butenyl)-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 15 2-(1E-Butenyl)-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.

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- 2-Propyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 2-Propyl-4-chloro-1[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.
- 2-Butyl-4-chloro-1-[2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-imidzole-5-carboxaldehyde.
- 2-(1E-Butenyl)-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-hydroxymethyl)imidazole.
- 25 2(1E-Butenyl)-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxaldehyde.
 - 2-Butyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)-biphenyl-4-yl)methyl]-imidazole-5-carboxylic acid.
- 2-Propyl-4-chloro-1-[(2-'(1H-tetrazol-5-yl)-biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.
 - 2-Propyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.
 - 2-Propyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxylmethyl)imidazole.
 - 35 2-Butyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-

yl)methyl]imidazole-5-carboxylic acid.

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- 2-Propyl-4-trifluoromethyl-1-[(2'-(carboxybiphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.
- 2-Propyl-4-pentafluoroethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 2-Propyl-1-[(2-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-4,5,-dicarboxylic acid.
- 2-Propyl-4-pentafluoroethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.
- 2-Propyl-4-pentafluoroethyl-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxaldehyde,

or its pharmaceutically acceptable salt thereof.

A more preferred embodiment of the invention is the method as recited above, using the imidazole angiotensin II receptor antagonist of formula I:

- 2-Butyl-4-chloro-1-[(2'-tetrazol-5-yl)biphenyl-4-yl]methyl]-5-(hydroxy-methyl)imidazole (also known as losartan); and
- 2-Butyl-4-chloro-1-[(2'-tetrazol-5-yl)biphenyl-4-yl]methylimidazole-5-carboxylic acid (also known as EXP-3174) or its pharmaceutically acceptable salt thereof. The most preferred embodiment is the method, as recited above, wherein the symptomatic heart failure patient is age 65

or older.

This invention also relates to a method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a

cardiac death by administering to a symptomatic heart failure patient a
therapeutically effective amount of an angiotensin II antagonist.

An embodiment of this invention is the method for

reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458, SL910102, UP2696, YM358, EMD66397, ME3221,

35 TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188,

EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498, and YM31472. The preferred angiotensin II receptor antagonists useful in this method are: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

Yet another embodiment of the invention is a method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

$$R^{6}$$
 R^{6}
 R^{6}
 R^{8}
 CH_{2}
 R^{1}
 R^{2}
 R^{3}

wherein:

15 \mathbb{R}^1 is:

$$\label{eq:conhomo} \begin{array}{c} {\rm CO_2H} \\ \text{4-CONHNHSO}_2{\rm CF}_3; \text{4-CONH-CHCH}_2{\rm C}_6{\rm H}_5 \end{array}; \\ \text{(I-isomer)} \end{array}$$

$$R^{13}$$
 R^{13} R^{13}

$$\begin{array}{c} O \\ -\ddot{\mathbb{C}} - \mathrm{NHSO_2} - \cdot \cdot (\mathrm{CH_2})_{\mathrm{s}} \end{array} ;$$

R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃;

R3 is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl - 24 -

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or cycloalkylalkynyl 5 to 10 carbon atoms; $(CH_2)_s Z(CH_2)_m R^5$ optionally substituted with F or CO_2R^{14} ; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

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R⁷ is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where v=1-6, C₆F₅; CN;

straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

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R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH₂)_m-imidazol-1-yl; -(CH₂)_m-1,2,3-triazolyl optionally substituted with one or two group selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms; -(CH₂)_s tetrazolyl;

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-(CH₂)_{n-1}CH-R¹¹; -(CH₂)_nOCR¹⁴; -(CH₂)_nSR¹⁵; OR¹⁷

25

R14 O O -CH=CH(CH2)sCHOR15; -CH=CH(CH2)sCR16; -CR16;

-CH=

-CH=CH(CH₂)_sOCR¹¹; (CH₂)_s-CH-COR¹⁶; CH₃

30

O Y Y Y (CH₂)_nCR₁₆; -(CH₂)_nOCNHR₁₀; -(CH₂)_nNR₁₁COR₁₀;

5

O -(CH₂)_nNR¹¹CNHR¹⁰; -(CH₂)_nNR¹¹SO₂R¹⁰; Y -(CH₂)_nNR¹¹CR¹⁰; -(CH₂)_mF; -(CH₂)_mONO₂; -CH₂N₃; -(CH₂)_mNO₂; -CH=N-NR¹¹R¹⁷;

-(CH₂)_m-N ; -(CH₂)_s N=NR⁴

$$-(CH_2)_s$$
 $N-N$
 CF_3 ; $-(CH_2)_n-N$
 CH_3O

$$-CH=N-NH-SO_2$$
; or $-CH=N-NH-N$

R²⁴ O R⁹ is: -CH-OCR²¹;

10

R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)pC6H5;

15 R11 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

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R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

-PO₃H₂; -C(CF₃)₂OH; -NHSO₂CH₃; -NHSO₂CF₃;

-NHCOCF3; -CONHOR12; -SO2NH2;

—CONHNHSO₂CF₃;
$$N-N$$
 or $N=N$ NH R⁴

R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;

R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;

R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R¹⁸ and R¹⁹ independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 $(CH_2)_t$ Q ;

25

Q is NR20, O or CH2;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or - CHCH2CO2CH3;
NH2

R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as (CH2)_u, where u is 3-6;

10 R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

—
$$NHSO_2$$
— CH_3 ; or — $NHSO_2$ — CH_3

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R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;

R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;

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R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;

R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;

X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23

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-NHC(R27)(R28)-, -NR23SO2-, -SO2NR23-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH2CH2-, -C(R27)(R28)NH-,

Y is O or S;

Z is O, NR11, or S;

5 m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

q is 2 to 3;

r is 0 to 2;

10 s is 0 to 5;

15

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds; provided that:

- (1) the R¹ group is not in the ortho position;
- (2) when R¹ is

$$-x$$
 R^{13}
 R^{2}

20 X is a single bond, and R13 is CO₂H, or

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

5 (3) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

- (4) when R1 is 4-CO₂H or a salt thereof, R6 cannot be S-alkyl;
- when R1 is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
 - (6) when R1 is

$$-X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

20

X is -OCH₂-, and R¹³ is 2-CO₂H, and R⁷ is H then R⁶ is not C₂H₅S;

(7) when R^1 is

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and R6 is n-hexyl then R7 and R8 are not both hydrogen;

10 (8) when R^1 is

R6 is not methoxybenzyl;

15

- (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
- (10) when r=0, R^1 is

$$X = \begin{pmatrix} R^{13} \\ - R^3 \end{pmatrix}$$

X is -NH-C-, R13 is 2-NHSO₂CF₃, and R6 is n-propyl, then R7 and R8 are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^2 \end{array} \right),$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

$$R^1 = X - \begin{pmatrix} R^{13} \\ R^2 \end{pmatrix}$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ | = \\ R^2 \end{array} \right)$$

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X is a single bond, R7 is Cl, and R8 is -CHO, then R13 is not 4-(tetrazol-5-yl).

A preferred embodiment of the method as recited above, wherein the angiotensin II receptor antagonist is losartan or EXP-3174 or a pharmaceutically acceptable salt thereof. The most preferred embodiment is the method, as recited above, wherein the symptomatic heart failure patient is age 65 or older.

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This invention also concerns a method for reducing mortality and sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist.

An embodiment of this invention is the method for reducing mortality and sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458, SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332,

CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498, and YM31472. The preferred angiotensin II receptor antagonists useful in this method are: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-

Yet another embodiment of the invention is a method for reducing mortality and sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist formula I

biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

wherein:

R¹ is:

 CO_2H 4-CONHNHSO $_2CF_3$; 4-CONH—CHCH $_2C_6H_5$; (*I*-isomer)

$$HO_2C$$
 R^{11}
 $A-CON$
 HO_2C
 R^{11}
 $A-N$
 R^{11}
 $A-N$
 R^{11}
 $A-N$
 R^{11}
 $A-N$
 R^{11}
 $A-N$
 R^{11}
 $A-N$
 R^{11}
 R^{1

$$A = N$$
 $A = N$
 $A =$

$$\begin{array}{c} {\rm O} \\ {\rm -C-NHSO_2---} \ ({\rm CH_2})_{\rm s} \end{array} ;$$

R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃;

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

R⁵ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl

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or cycloalkylalkynyl 5 to 10 carbon atoms; $(CH_2)_s Z(CH_2)_m R^5$ optionally substituted with F or CO₂R14; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

5

R7 is H, F, Cl, Br, I, NO2, $C_{\nu}F_{2\nu+1}$, where v=1-6, C6F5; CN;

straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

10

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R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; $-(CH_2)_m$ -imidazol-1-yl; $-(CH_2)_m$ -1,2,3-triazolyl optionally substituted with one or two group selected from CO2CH3 or alkyl of 1 to 4 carbon atoms; -(CH2)s

20

tetrazolyl;

-(CH₂)_{n-1}CH-R¹¹; -(CH₂)_nOCR¹⁴; -(CH₂)_nSR¹⁵; OR¹⁷

25

R14 O O -CH=CH(CH2)sCHOR15; -CH=CH(CH2)sCR16; -CR16;

30

-CH=CH(CH₂)_sOCR¹¹; (CH₂)_s-CH-COR¹⁶;

O Y Y Y Y (CH2)nCR16; -(CH2)nOCNHR10; -(CH2)nNR11COR10;

-(CH₂)_nNR¹¹CNHR¹⁰; -(CH₂)_nNR¹¹SO₂R¹⁰;

-(CH₂)_nNR₁₁CR₁₀; -(CH₂)_mF; -(CH₂)_mONO₂; -CH₂N₃; -(CH₂)_mNO₂; -CH=N-NR₁₁R₁₇;

$$-(CH2)m-N ; -(CH2)s N=N R4$$

$$-(CH2)s - N N CF3; - (CH2)n - N CH3O$$

$$-- (CH2)n-1C-N N- CH3O;$$

R²⁴ Q R⁹ is: -CH-OCR²1;

5

- 10 R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)_pC6H5;
- R¹¹ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

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R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

-PO3H2; -C(CF3)2OH; -NHSO2CH3; -NHSO2CF3;

-NHCOCF3; -CONHOR12; -SO2NH2;

- R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
 - R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
 - R16 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;
- R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
 - R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 $Q;$

25

Q is NR20, O or CH2;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or - CHCH2CO2CH3; NH2

R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

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R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

15

- R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl:
- R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;

20

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- R²⁹ and R³⁰ are independently alkyl of 1-4 carbon atoms or taken together are -(CH₂) $_q$ -;
- R³¹ is H, alkyl or 1 to 4 carbon atoms, -CH₂CH=CH₂ or -CH₂C₆H₄R³²;

X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23

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-NHC(R27)(R28)-, -NR23SO2-, -SO2NR23-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH2CH2-, -C(R27)(R28)NH-,

Y is O or S;

Z is O, NR11, or S;

5 m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

q is 2 to 3;

r is 0 to 2;

10 s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds; provided that:

- (1) the R1 group is not in the ortho position;
- 15 (2) when R¹ is

$$-X - \left(\begin{array}{c} R^{13} \\ - \\ R^{2} \end{array} \right)$$

20 X is a single bond, and R13 is CO₂H, or

then R^{13} must be in the ortho or meta position; or when R^1 and X are as above and R^{13} is NHSO₂CF₃ or NHSO₂CH₃, R^{13} must be ortho;

5

(3) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ - \\ R^{2} \end{array} \right)$$

10

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

- (4) when R1 is 4-CO2H or a salt thereof, R6 cannot be S-alkyl;
- when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
 - (6) when R^1 is

$$-x - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

20

X is -OCH₂-, and R^{13} is 2-CO₂H, and R^{7} is H then R^{6} is not C₂H₅S;

(7) when R^1 is

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and R6 is n-hexyl then R7 and R8 are not both hydrogen;

10 (8) when R^1 is

R6 is not methoxybenzyl;

15

- (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
- (10) when r=0, R^1 is

$$X = \begin{bmatrix} R^{13} \\ - R^{3} \end{bmatrix},$$

20

X is -NH-C-, R¹³ is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ = \end{array} \right) = R^3$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ = \\ R^2 \end{array} \right)$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \begin{pmatrix} & & & \\$$

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X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 4-(tetrazol-5-yl).

A preferred embodiment of the invention is the method as recited above, wherein the angiotensin II receptor antagonist is losartan or EXP-3174 or a pharmaceutically acceptable salt thereof. The most preferred embodiment is the method, as recited above, wherein the symptomatic heart failure patient is age 65 or older.

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Additionally, the invention concerns a method for preventing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist.

An embodiment of this invention is the method for preventing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan,

- valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458, SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188,
- EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156,
 LRB057, LY266099, LY301875, PD123177, PD126055, SC51757,
 SC54629, U96849, UK77778, WAY126227, WK1260, WK1492,
 YH1498, andYM31472. The preferred angiotensin II receptor
 antagonists useful in this method are: candesartan cilexetil, eprosartan,
- irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

Yet another embodiment of the invention is a method for preventing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

$$R^6$$
 R^7
 R^8
 $(CH_2)_r$
 R^1
 R^3

5

wherein:

 R^1 is:

R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃;

CONHOR¹²;
$$SO_2NH_2$$
; $N-N$
N; aryl; or furyl;

5

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

10

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl or cycloalkylalkynyl 5 to 10 carbon atoms; (CH₂)_SZ(CH₂)_mR⁵ optionally substituted with F or

20

CO₂R¹⁴; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

 R^7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where $\nu=1-6$, C₆F₅; CN;

25

O—C-R¹⁶; straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH₃, CF₃, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

30

	R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH ₂) _m -imidazol-1-yl; -(CH ₂) _m -1,2,3-triazolyl
5	optionally substituted with one or two group selected from CO2CH3 or alkyl of 1 to 4 carbon atoms; -(CH2)s
	tetrazolyl;
10	O -(CH ₂) _{n-1} CH-R ¹¹ ; -(CH ₂) _n OCR ¹⁴ ; -(CH ₂) _n SR ¹⁵ ; OR ¹⁷
	R14 O O -CH=CH(CH2)sCHOR15; -CH=CH(CH2)sCR16; -CR16;
15	O -CH=CH(CH ₂) _s OCR ¹¹ ; (CH ₂) _s -CH-COR ¹⁶ ; CH ₃
	O Y Y -(CH ₂) _n CR ₁₆ ; -(CH ₂) _n OCNHR ₁₀ ; -(CH ₂) _n NR ₁₁ COR ₁₀ ;
	O -(CH ₂) _n NR ¹¹ CNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ SO ₂ R ¹⁰ ;
20	Y -(CH ₂) _n NR ₁₁ CR ₁₀ ; -(CH ₂) _m F; -(CH ₂) _m ONO ₂ ; -CH ₂ N ₃ ; -(CH ₂) _m NO ₂ ; -CH=N-NR ₁₁ R ₁₇ ;

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow NH ;$$

$$-(CH_{2})_$$

R²⁴ O R⁹ is: -CH-OCR21;

- 5 R¹⁰ is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH₂)_pC₆H₅;
- R¹¹ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

15 O O O OH O

-O-S-OH; -O-P-OH; -SO₃H; -NHP-OH; -C - P - OH

OH OH OH OH R²⁷OH

-PO₃H₂; -C(CF₃)₂OH; -NHSO₂CH₃; -NHSO₂CF₃;
-NHCOCF₃; -CONHOR¹²; -SO₂NH₂;

$$\begin{array}{c}
N-N \\
N \\
N
\end{array}$$

$$\begin{array}{c}
N-N \\
N \\
N
\end{array}$$

$$\begin{array}{c}
N-N \\
N
\end{array}$$

- R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
- 10 R16 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH2)_pC6H5, OR17, or NR18R19;
 - R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- 15
 R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

20 O is NR²⁰, O or CH₂;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

25 R21 is alkyl of 1 to 6 carbon atoms, -NR²²R²³, or - CHCH₂CO₂CH₃;
NH₂

- R^{22} and R^{23} independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;
- 5 R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

- 10 R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
 - R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;
- 15 R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;
 - R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;

20
X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23

-NHC(R27)(R28)-, -NR23SO₂-, -SO₂NR23-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH₂CH₂-, -C(R27)(R28)NH-,

Y is O or S;

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Z is O, NR11, or S;

m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

5 q is 2 to 3;

r is 0 to 2;

s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds;

10 provided that:

- (1) the R¹ group is not in the ortho position;
- (2) when R^1 is

$$-x - \left(\frac{R^{13}}{R^2} \right)$$

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X is a single bond, and R13 is CO₂H, or

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then R13 must be in the ortho or meta position; or when R1 and X are as above and R13 is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

(3) when R1 is

$$-X - \left(\frac{R^{13}}{R^2} \right)$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

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- (4) when R¹ is 4-CO₂H or a salt thereof, R⁶ cannot be S-alkyl;
- (5) when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
 - (6) when R¹ is

$$-x$$
 R^{13}
 R^{2}

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X is -OCH2-, and R^{13} is 2-CO2H, and R^7 is H then R^6 is not C2H5S;

(7) when R^1 is

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and R6 is n-hexyl then R7 and R8 are not both hydrogen;

(8) when R^1 is

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R6 is not methoxybenzyl;

- 10 (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
 - (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ | = R^3 \end{array} \right)$$

X is -NH-C-, R¹³ is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^2 \end{array} \right),$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO₂CH₃;

(12) when r=1,

$$R^1 = X - \begin{cases} R^{13} \\ - R^3 \end{cases}$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^{1}=X$$
 R^{2}
 R^{13}
 R^{2}

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X is a single bond, R^7 is Cl, and R^8 is -CHO, then R^{13} is not 4-(tetrazol-5-yl).

A preferred embodiment of the invention is the method as recited above, wherein the angiotensin II receptor antagonist is losartan or EXP-3174 or a pharmaceutically acceptable salt thereof. The most preferred embodiment is the method, as recited above, wherein the symptomatic heart failure patient is age 65 or older.

EXAMPLE 1

ELITE Study

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Patients and methods

The ELITE study was a prospective double-blind, randomized, parallel, captopril-controlled clinical trial conducted at 125 centers in the United States, Europe and South America. The study was approved by institutional review boards at each site; all patients gave written informed consent. An independent Data and Safety Monitoring Committee monitored the progress of the study. See Pitt B, Chang P, Timmermans P. Angiotensin II receptor antagonists in heart failure: Rationale and design of the Evaluation of Losartan in the Elderly (ELITE) Trial. Cardiovascular Drugs and Therapy 1995; 9: 693-700.

20 Patient Population

Patients were 65 years or older (two thirds 70 years or older) with symptomatic heart failure (New York Heart Association Class II-IV), decreased left ventricular ejection fraction of 40 percent or less, and had no history of prior ACE inhibitor therapy. A detailed description of enrollment and exclusion criteria and study design has been presented previously. See Pitt B, Chang P, Timmermans P. Angiotensin II receptor antagonists in heart failure: Rationale and design of the Evaluation of Losartan in the Elderly (ELITE) Trial. Cardiovascular Drugs and Therapy 1995; 9: 693-700.

Randomization and Study Therapy

Following a two-week placebo run-in, patients were randomized to 48 weeks of active therapy, either to captopril 6.25 mg titrated to 12.5 mg, 25 mg and then 50 mg three times daily (plus placebo for losartan) or to losartan 12.5 mg, titrated to 25 mg and then

50 mg once daily (plus placebo for captopril). Titration generally occurred at 7-day intervals as tolerated. Treatment with all other concomitant cardiovascular therapies was permitted with the exception of open-label ACE inhibitors. Randomized patients were stratified by age (less than 70 and 70 years or greater).

Evaluation of Patients

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Clinical assessments were performed weekly during titration and at three-month intervals thereafter. Laboratory evaluations were performed at weeks three, six, and twelve, and at three month intervals thereafter.

Study Endpoints

The primary endpoint of the study was a safety measure of 15 the development of renal dysfunction, defined as an increase in serum creatinine >0.3 mg/dL from baseline that was confirmed by a repeat measurement in 5 to 14 days of the first determination, during continued treatment. All-cause mortality and heart failure hospitalizations were each prespecified endpoints, and the composite of 20 death and/or heart failure hospitalizations was added as the secondary endpoint by protocol amendment based on data from two placebocontrolled 12 week exercise studies (of approximately 350 patients each), demonstrating a beneficial effect on this endpoint. See Example 2) Hypotension-related symptoms, clinically important serum potassium 25 increases (≥ 0.5 mEq/L), and cough, all originally secondary endpoints, were moved to tertiary endpoints. All deaths (including cause of death) and hospitalizations were adjudicated by an independent Clinical Endpoint Adjudication Committee, blinded to study treatment (see panel for mortality classification). Other prospectively-defined measures 30 included myocardial infarction/hospitalization for unstable angina, worsening of heart failure, New York Heart Association functional classification, discontinuation from the study due to study drug intolerance, changes in neurohormonal profile.

35 Panel: Mortality Classification

• Sudden Cardiac Death: Death occurring without warning or within 1 hour of symptoms

• <u>Death due to Progressive Heart Failure</u>: Death preceded by worsening signs and/or symptoms of heart failure, including cardiogenic shock

• <u>Fatal Myocardial Infarction</u>: Death associated with autopsy-verified myocardial infarction or death within 28 days of a hospital-verified acute myocardial infarction provided no other cardiac or non-cardiac cause of death is found

• <u>Death due to Other Cardiac Causes</u>: Death due to other cardiac causes, such as arrhythmia

• <u>Death due to Other Vascular Causes</u>: Death due to vascular events such as stroke, pulmonary embolus, ruptured aneurysm, etc.

• Death due to non-cardiovascular causes: Death not due to any cardiac or vascular events

Statistical Methods

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Drugs and Therapy 1995; 9: 693-700. Analysis of renal dysfunction (increases in serum creatinine) was based upon a modified intent-to-treat population; i.e., all patients were analyzed according to their randomization group, and an endpoint was declared only if initial and confirmatory elevations occurred while on double-blind therapy. Patients who discontinued from the study without meeting this endpoint were censored in the time-to-event analysis at the time of study discontinuation.

Analyses of deaths and heart failure hospitalizations (adjudicated endpoints) were based on an intent-to-treat population; all patients discontinued prematurely were followed through the specified 48-week period of the study. Patients not meeting the endpoint were censored in the time-to-event analysis either at the time of study completion (for patients who completed) or at the end of the 48-week follow-up period (for patients who discontinued).

For all time-to-event data, survival analyses were based upon the log-rank test. The effect of treatment group in the model was tested controlling for the stratification factor (age category [less than 70 or 70 years and older]). The time to *first* event was used for each endpoint. Risk reductions were based upon Mantel-Haenszel adjusted (for age catagory) relative risk estimates.

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The primary endpoint was tested at the 5% level of significance. No multiplicity adjustments were made for the test of the secondary endpoint or for other tests of interest; unadjusted p-values are presented.

Over approximately a two-year recruitment period, 722 patients enrolled, 352 were randomized to losartan and 370 to captopril (Figure 1). The two treatment groups were similar with respect to all baseline characteristics (Table 1). Concomitant therapies during the study were similar between the two treatment groups; diuretics were used in over 70 percent of patients, digitalis in over 55% of patients and non-ACE inhibitor vasodilatory drugs (including hydralazine and nitrates) in over 50 percent of patients in both treatment groups. Three hundred patients (85 percent) were titrated to the target dose of losartan 50mg once daily, and 321 patients (84 percent) in the captopril group were titrated to the target dose of 50mg three times a day. Seventy-five percent of losartan-treated patients remained on the targeted dose of 50mg daily (mean daily dose 42.6 mg per day), and 71 percent of captopril patients remained on the targeted dose of 50mg three times a day (mean daily dose 122.7 mg per day).

Table 1. Baseline Clinical Characteristics and Drug Therapy, According to

Tı	reatment Group				
		LOS	SARTAN	CAP	OPRIL
	CHARACTERISTIC	(1	n=352)	(n=	=370)
Sex		nun	nber (perce	ent) of p	atients
	Male	234	(66.5)	248	(67.0)
	Female	118	(33.5)	122	(33.0)
Age (y	years)				
	<70	95	(27.0)	119	(32.2)
	≥70		(73.0)	251	(67.8)
	Mean (S.D.)	74	(5.8)	73	(6.1)
Race					
	Caucasian	320	(90.9)	326	(88.1)
	Black	16	(4.6)	18	(4.9)
	Other	16	(4.6)	26	(`7.0)
Etiolo	gy of Heart Failure †				` ,
Enoio	Ischemic Heart Disease	242	(68.8)	250	(67.6)
	Non-Ischemic Heart Disease	110	(31.3)	120	(32.4)
NYH	A Functional Class ***	231	(65.6)	237	(64.1)
	Щ		(33.0)	126	(34.1)
	m	5	(1.4)	7	(1.9)
ъ,	IV	-	(,	·	()
Drug	Therapy	260	(72.0)	275	(74.2)
	Diuretics Digitalia		(73.9) (56.5)	275	(74.3)
	Digitalis Hydralazine	129	(56.5) (3.4)	209 12	(56.5) (3.2)
	Nitrates		(51.1)	191	(51.6)
	Calcium Channel Blockers		(34.9)	121	(32.7)
	Potassium Supplement	91	(25.9)	89	(24.1)
	Anticoagulants	60	(17.0)	69	(18.6)
	Beta-Blockers	55	(15.6)	63	(17.0)
	Antiarrhythmics	37	(10.5)	39	(10.5)
_		•	(200)		(-0.0)
Secon	dary Diagnoses	104	(50.0)	100	(45.0)
	Myocardiai infarction		(52.3)	177	(47.8)
	Hypertension	201	(57.1)	212	(57.3)
	Atrial Fibrillation	86	(24.4)	82	(22.2)
	Diabetes mellitus	94 21	(26.7)	89	(24.1)
	Renal Insufficiency	32	(6.0) (9.1)	26 37	(7.0)
	Stroke Current smoker (cigarettes)	39		45	(10.0) (12.2)
	Current shloker (eigarettes)	37	(11.1)	(S.D.)	(12.2)
Fiecti	on Fraction (%)	31	(7.2)	30	(7.6)
			(0.4)	1.2	(0.4)
Serum	Creatinine (mg/dl)				•
Semm	Potassium (mEq/L) †††	4.3	(0.4)	4.3	(0.5)
Heart	Rate (beats per minute)	73	(11.7)	74	(10.4)
	Pressure		\ <i>\</i>	- •	(/
	Systolic (mmHg)	137	(17.6)	137	(19.1)
	Diastolic (mmHg)	79	(9.4)	79	(10.6)
Weigh	· ·	76	(33.1)	74	(33.8)

Key to Table 1:

- † Based on patient history.
- 11 NYHA denotes New York Heart Association.
- To convert to micromoles per liter, multiply by 88.4; mEq/L is the equivalent of millimoles per liter.

Renal Dysfunction

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There was no significant difference between losartan and captopril in terms of persistent increases (i.e., confirmed by a repeat measurement during continued treatment) in serum creatinine ≥0.3 mg/dL from baseline (10.5 percent versus 10.5 percent, p=0.629; Table 2). Single rises of serum creatinine ≥0.3 mg/dL were documented in 92 losartan-treated patients (26.1 percent) and 110 captopril-treated patients (29.7 percent), with a trend towards fewer events occuring in losartan-treated patients (14 percent risk reduction, 95 percent confidence intervals -0.09 to 0.32; p=0.060). Of these patients with single rises of serum creatinine ≥0.3 mg/dL, 68 percent had confirmation measurements performed while on active therapy per protocol; 55 percent of the patients who had these confirmation measurements met the endpoint.

Death and/or Heart Failure Hospitalizations

Follow-up data on death and hospitalizations were complete except for one losartan-treated patient who was discontinued after one dose of study medication. During the course of the study, death and/or 25 heart failure hospitalizations occurred in 33 of 352 losartan-treated patients (9.4 percent) compared to 49 of 370 captopril-treated patients (13.2 percent) (p=0.075). This decrease in death and/or heart failure hospitalizations observed with losartan versus captopril was entirely due 30 to a 46 percent decrease in total mortality (17 versus 32 patients [p=0.035]; Table 3). The cumulative survival curves (intent-to-treat) separated early and remained so throughout the 48-week study period (Figure 2). The observed decrease in total mortality was primarily due to a reduction in sudden cardiac death (five versus 14 patients, 35 p=0.043); only one patient died of progressive heart failure in each treatment group (Table 3). Fatal myocardial infarction occurred in one

losartan-treated patient versus four in the captopril arm. The effect of losartan versus captopril on mortality was generally a consistent observation across the different subgroups (Figure 3), except in a subpopulation of female patients (9 of 118 losartan-treated versus 8 of 122 captopril-treated female patients died).

Fewer losartan-treated patients were hospitalized overall for any reason during the 48-week study observation period than captopril-treated patients (86 [24.4 percent] versus 119 [32.2 percent]; p=0.018). The incidence of heart failure hospitalizations however was not different between treatment groups (both 5.7 percent) (Table 3).

Functional Status and Norepinephrine Levels

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VSDOCID- JWO GRROSTEAT I

New York Heart Association functional class improved similarly with losartan and captopril treatment (p≤0.001 versus baseline for each treatment group); 80 percent of losartan-treated patients and 82 percent of captopril-treated patients were classified as New York Heart Association Class I or II at the end of the study versus 66 percent of losartan-treated patients and 64 percent of captopril-treated patients at baseline (Figure 4). With regard to plasma norepinephrine levels, a three percent decrease from a geometric mean baseline of 469 pg/mL was observed at 48 weeks with losartan treatment, as compared to a five percent increase from a geometric mean baseline of 424 pg/mL with captopril (p = NS between treatment groups).

Table 2. Frequency of Increases in Serum Creatinine in All Patients and by Age

		P-VALUE ##			0.629		•			
	ISK REDUCTION	(95% CI) ^{††}			(-0.51, 0.36)		(-1.44, 0.59)		(-0.59, 0.40)	
	RISK	5)			0.02		0.00		0.05	
9-1		CRUDE EVENT RATE	number (percent) of	tients	(10.5)	(10.5)	(8.4)			
		CRUDEE	number	Ba	37	39	∞	10	53	29
		Z							257	
		TREATMENT			Losartan	Captopril	Losartan	Captopril	Losartan	Captopril
Table 2. I todacile) of the compa	AGE	CATEGORY [†] T			Overall		<70 years		≥70 years	•
1 acre 2: 1 100		EVENT			>0.3 ma/d1	Local Inglan	Increase (gonfirmed	Commission	Within 5 14 Days)	J-14 L/4ys)

Treatment comparisons were performed for overall group only.

†† Risk Reduction is the reduced risk of experiencing the related endpoint on losartan compared to captopril (a negative number denotes an increase in risk); overall estimates control for age category; CI denotes confidence interval.

111 p-value based upon log-rank test (survival analysis) with age category included as a stratification factor in the model.

Table 3. Mortality/Heart Failure Hospitalizations and Cause of Death

VARIABLE	LOS	SARTAN n=352	CAF	COSARTAN CAPTOPRIL n=352 n=370	RISK	RISK REDUCTION	P-VALIE #
:	umu	number (percent) of patients	ent) of	patients			
Death and/or Hospitalization for Heart Failure	33	(9.4)	49	(13.2)	0.32	0.32 (- 0.04, 0.55)	0.075
Total Mortality	17	(4.8)	32	(8.7)	0.46	0.46 (0.05, 0.69)	0.035
Sudden Death Progressive Heart	2	(1.4)	4 1	(3.8)	0.64	0.64 (0.03, 0.86) -0.11 (-20.23, 0.94)	0.043 0.930
Failure Fatal Myocardial	-	(0.3)	4	(1.1)	0.76	0.76 (- 0.83, 0.97)	0.172
infaction Other Vascular Non-cardiovascular	N N	(1.4)	√ 0 ∞	(1.4)	-0.03 0.35	-0.03 (- 2.63, 0.71) 0.35 (- 0.94, 0.78)	0.958 0.434
Hospitalization for Heart Failure	20	(5.7)	21	(5.7)	0.04	0.04 (- 0.74, 0.47)	0.894

Risk Reduction is the reduced risk of the related endpoint on losartan compared to captopril (a negative number denotes an increase in risk); estimates control for age category; CI denotes confidence interval.

P-value based upon log-rank test (survival analysis) with age category included as a stratification factor in the model.

ELITE is the first long-term (48 weeks) study to compare the effect of treatment with losartan, an angiotensin II type 1 receptor antagonist, to an ACE inhibitor (captopril) in patients with symptomatic heart failure and systolic left ventricular dysfunction. Captopril was chosen as the comparative ACE inhibitor in this study because it had been suggested to have less adverse renal effects than longer-acting ACE inhibitors. (See Packer, M., Lee, W.H., Yushak, M., Medina, N.: Comparison of Captopril and Enalapril in patients with severe chronic heart failure. N Engl J Med; 315: 847-853.) The incidence of persistent renal dysfunction, as defined by a persisting increase in serum creatinine of 0.3 mg/dL (the primary endpoint), was not different between losartan- and captopril-treated patients (both 10.5%). The observation overall that both losartan and captopril were relatively well tolerated with regard to renal function was evident, given that less than 2% of patients discontinued for this reason in either group. This is of clinical relevance for treating older patients with heart failure.

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The results of this study demonstrate that treatment with losartan resulted in a 46% reduction in all-cause mortality compared to captopril, a drug with demonstrated survival benefit in several studies. 20 (See Pfeffer MA, Braunwald E, Moye LA, et al. on behalf of the SAVE Investigators. Effect of captopril on mortality and morbidity in patients with left ventricular dysfunction after myocardial infarction. Results of the Survival and Ventricular Enlargement Trial. N Engl J Med 1992; 327: 669-677; Fonarow GC, Chelimsky-Fallick C, Warner Stevenson L, 25 et al. Effect of direct vasodilation with hydralazine versus angiotensinconverting enzyme inhibition with captopril on mortality in advanced heart failure: the Hy-C trial. J Am Coll Cardiol 1992; 19: 842-850; and ISIS Collaborative Group OU. ISIS-4: Randomized study of oral isosorbide mononitrate in over 50,000 patients with suspected acute 30 myocardial infarction. Circulation 1993; 88: I394.) The cumulative survival benefit of losartan was observed early during the study, persisted throughout the 48-week treatment period, and was consistent among all subgroups except in female patients. It should be noted that the number of female patients enrolled in this study was relatively small; 35 and the mortality benefit could not be demonstrated in this study. The

greater drop-out rate in the captopril-treated patients did not account for the beneficial effects of losartan on total mortality; the treatment difference in total mortality was primarily due to those who remained on active therapy.

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An improvement in survival with losartan of similar magnitude has also been observed in two placebo-controlled 12-week exercise studies. See Klinger G, Jaramillo N, Ikram H, et al. Effects of losartan on exercise capacity, morbidity and mortality in patients with symptomatic heart failure. J Am Coll Cardiol 1996 (in press); (Abstract). The three-month control group mortality rates in the exercise studies and the present study were comparable to the placebo and enalapril mortality rates respectively in the Studies of Left Ventricular Dysfunction (SOLVD) trial (Figure 5). See The SOLVD Investigators. Effect of enalapril on survival in patients with reduced left ventricular ejection fractions and congestive heart failure. N Engl J Med 1991; 325: 293-302. Despite the limitations of cross-study comparisons, treatment with losartan in the exercise studies and the present study was associated with comparably low mortality rates which were less than the observed mortality rates for both placebo and enalapril in SOLVD.

Prior to the use of ACE inhibitors, progressive heart failure accounted for approximately 50% of deaths due to heart failure. See Applefield MM. Chronic congestive heart failure: Where have we been? Where are we heading? Am J Med 1986; 80: 73-77. In patients with mild to moderate heart failure treated with an ACE inhibitor, death due to progressive heart failure has diminished so that sudden cardiac death is currently the predominant mode of death. For example, in the recently discontinued Survival With Oral d-Sotalol (SWORD) trial, in which patients with mild to moderate heart failure or left ventricular dysfunction were randomized to d-Sotolol or placebo on a background of usual therapy including an ACE inhibitor, arrhythmic death accounted for approximately two-thirds of total mortality in the placebo group, while progressive heart failure for only one-sixth. See Waldo AL, Camm AJ, deRuyter H, et al. for the SWORD Investigators. Effect of d-sotalol on mortality in patients with left ventricular dysfunction

after recent and remote myocardial infarction. Lancet 1996; 348: 7-12. In the present study, sudden cardiac death was also the most prominent cause of death in the captopril-treated patients. Of interest, the observed reduction in mortality on losartan compared to captopril-treated patients in the present study was primarily due to a reduction in sudden cardiac death (see Table 3). Few patients died due to progressive heart failure or to fatal myocardial infarction in either treatment group, although numerically, fatal myocardial infarction deaths occurred less with losartan.

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10 The mechanism by which losartan reduces sudden cardiac death in comparison to captopril is uncertain. Whether the reduction in sudden cardiac death was due to more complete blockade of angiotensin II effects or an intrinsic antiarrhythmic/antifibrillatory property of losartan remains to be determined. ACE activity may not be completely 15 suppressed by the presently studied captopril dosing regimen (target dose of 50 mg three times daily), which is a regimen considered to have a mortality benefit. See Cohn JN. The management of chronic heart failure. N Engl J Med 1996; 335: 490-498. In addition angiotensin II may also be formed by non-ACE-dependent pathways. (See Miura S, 20 Ideishi M, Sakai T, et al. Angiotensin II formation by an alternative pathway during exercise in humans. J Hypertension 1994; 12: 1177-1181; Urata H, Kinoshita A, Misono KS, Bumpus FM, Husain A. Identification of a highly specific chymase as the major angiotensin IIforming enzyme in the human chymase. J Biol Chem 1990; 265: 22348-25 22357; Urata H, Strobel F, Ganten D. Widespread tissue distribution of human chymase. J Hypertension 1994; 12: S17-S22; and Aldigier JC, Huang H, Dalmay F, et al. Angiotensin-converting enzyme inhibition does not suppress plasma angiotensin II increase during exercise in humans. J Cardiovasc Pharmacol 1993; 21: 289-295.) It is possible that 30 more complete blockade of angiotensin II effects by losartan may result in more complete suppression of catecholamines at the tissue level. See Brasch H, Sieroslawski L, Dominiak P. Angiotensin II increases norepinephrine release from atria by acting on angiotensin subtype 1 receptors. Hypertension 1993; 22: 699-704. Furthermore, bradykinin, 35 which is known to release norepinephrine, is not elevated with direct

angiotensin II blockade compared with ACE inhibitor therapy. (See Minisi AJ, Thames MD. Distribution of left ventricular sympathetic afferents demonstrated by reflex responses to transmural myocardial ischemia and to intracoronary and epicardial bradykinin. Circulation 1993; 87: 240-246; and Timmermans P, Wong PC, Chiu AT, et al. Angiotensin II receptors and angiotensin II receptor antagonists. Pharmacol Reviews 1993; 45: 205-251.) A small decrease in mean plasma norepinephrine was observed in the present study with losartan compared to captopril. Plasma catecholamine levels may not, however, reflect local cardiac tissue levels, and resting values may not reflect transient increases during stress or ischemia.

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In this study, heart failure status appeared to improve to a similar degree in both treatment groups. New York Heart Association functional class improved significantly and to a comparable extent from baseline following long-term treatment with both losartan and captopril (Figure 4). The rate of hospitalization for heart failure in patients treated with losartan was also similar to that observed in patients treated with captopril. Another similarity was the low incidence of death due to progressive heart failure (less than one percent for both losartan and captopril-treated patients), suggesting a similar beneficial effect for both treatments. Overall hospitalizations were however less frequent with losartan treatment versus captoril.

In conclusion, in older patients with symptomatic heart

failure, losartan was better tolerated compared to captopril, with fewer patients discontinuing therapy due to adverse effects or being hospitalized, but was not different from captopril in causing persistent renal dysfunction defined as a sustained increase in serum creatinine. The effects on progressive heart failure hospitalization and improvement in New York Heart Association functional class were similar in the two treatment groups. There was, however, a decrease in all-cause mortality observed with losartan compared to captopril treatment, mainly due to a reduction in sudden cardiac death. These findings suggest that treatment with losartan may offer important

therapeutic benefits compared to an ACE inhibitor, and may have far reaching implications for the treatment of patients with heart failure.

EXAMPLE 2

5 Exercise Study—Protocols A & B

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Protocols A & B were, multicenter, double-blind, randomized, parallel, placebo-controlled studies to investigate the effects of losartan on the exercise capacity and clinical status of patients over the age of 18 with symptomatic heart failure.

Approximately three hundred and fifty (350) patients with symptomatic heart failure (NYHA Functional Class II-IV, at least 50% Class III-IV) and left ventricular ejection fraction ≤40% were randomized to receive either losartan or placebo once daily for 12 weeks in each study. Following a minimum 15-day baseline placebo period, during which a minimum of 5 and maximum of 8 sets of baseline exercise tests (treadmill exercise test and 6-minute walk test) were performed, patients were randomized to treatment with losartan or placebo in a 2:1 ratio, respectively. According to the protocol's, patients must not have received an ACE-inhibitor for at least 6 weeks prior to the third baseline exercise test visit. Treatment with an ACE-inhibitor was not allowed at any time during these studies.

This report summarizes the results of the Meta-analysis performed (as proposed in the Data Analysis Plan) on the observed mortality and congestive heart failure (CHF) hospitalization across these two studies. The mortality and CHF hospitalization were not predefined endpoints in the protocols but it was intended that they be analyzed as a part of safety. Thus results from these analyses should not be interpreted as a test of significance as there were no predefined hypotheses corresponding to these safety parameters. However, these results can simply be regarded as a measure of the disparity between the losartan and placebo groups with respect to the observed results across the two studies. Odds ratio's with 95% confidence intervals have been provided for these variables.

Peto-modified Mantel-Haenszel (Fleiss, J.L., Statistical
Methods for Rates and Proportions, Wiley, NY, 1981.) method was used

to carry out the Meta-analysis of death and CHF hospitalization rates in the two 12 week exercise studies. This method assumes homogeneity across the studies and provides a test for heterogeneity. Results of the Meta-analysis are displayed in Tables 4-6.

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A total of three hundred and fifty one (351) patients were enrolled in the protocol A study; 237 patients were randomized to receive losartan and 114 patients were randomized to receive placebo. There were 4 (3.5%) deaths in the placebo group as compared to 4 (1.7%) in the losartan group during the double-blind therapy or as related to serious adverse events (AE) during double-blind. Additionally there were 10 (8.8%) CHF hospitalizations in the placebo group as compared to 10 (4.2%) in the losartan group during the double-blind therapy or as related to serious AE during double-blind. One patient included in the losartan group discontinued due to worsening HF and was hospitalized 2 days after discontinuation from double-blind period.

Table 4 provides the odds ratio for the Death and CHF hospitalization in the protocol A exercise study. The odds of dying in the placebo group are 2.1 times the odds of dying in the losartan group while the odds of CHF hospitalization in the placebo group are 2.2 times the odds of CHF hospitalization in the losartan group. However, since 1.0 is included in both the 95% confidence intervals the possibility that the odds of dying (or CHF hospitalization) in the placebo group is the same as odds of dying (or CHF hospitalization) in the losartan group can not be excluded (see Table 4).

A total of three hundred and eighty five (385) patients were enrolled in the protocol B exercise study; 254 patients were randomized to receive losartan and 131 patients were randomized to receive placebo. There were 9 (6.9%) deaths in the placebo group as compared to 3 (1.2%) in the losartan group during the double-blind therapy or as related to serious AE during double-blind. Additionally there were 7 (5.3%) CHF hospitalizations in the placebo group as compared to 5 (2.0%) in the losartan group during the double-blind therapy or as related to serious AE during double-blind.

Table 5 provides the odds ratio for the Death and CHF hospitalization in the protocol B exercise study. The odds of dying in the placebo group are 5.6 times the odds of dying in the losartan group while the odds of CHF hospitalization in the placebo group are 2.7 times the odds of CHF hospitalization in the losartan group. The 95% confidence interval for odds ratio of death was (1.61, 19.34). This implies that the odds of dying in the placebo group are greater than the odds of dying in the losartan group. However, since 1.0 is included in the 95% confidence interval for odds ratio of CHF hospitalization, the possibility that the odds of CHF hospitalization in the placebo group is same as the odds of CHF hospitalization in the losartan group can not be excluded (see Table 5).

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Since the results from protocols A and B are consistent, a safety Meta-analysis as proposed in the Data Analysis Plan has been performed. Homogeneity between the two studies with respect to odds ratio's for death and CHF hospitalization was found.

Table 6 provides the results of the Meta-analysis of the two 12 week exercise studies. The odds of dying in the placebo group are 3.5 times the odds of dying in the losartan group while the odds of CHF hospitalization in the placebo group are 2.4 times the odds of CHF hospitalization in the losartan group. The 95% confidence intervals for odds ratio's with respect to death and CHF hospitalization are (1.43, 8.77) and (1.19, 4.76), respectively. Thus the odds of dying in the placebo group are significantly greater than the odds of dying in the losartan group. Similarly the odds of CHF hospitalization in the placebo group are significantly greater than the odds of CHF hospitalization in the losartan group.

TABLE 4
CLINICAL EVENTS—PROTOCOL A

	PBO (N=131)	Losartan (N=254)	Odds Ratio	95% Confid Intervals Lower	ence Upper
Death ^b	3.5%	1.7%	2.1	0.56	7.96
CHF Hospitalization (during DB)	8.8%	4.2% a	2.2	0.90	5.28

b: during double-blind therapy or was related to SAE during double-blind

5 a: one patient discontinued due to worsening HF and was hospitalized 2 days after discontinuation from double-blind.

TABLE 5
CLINICAL EVENTS — PROTOCOL B

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	PBO (N=131)	Losartan (N=254)	Odds Ratio	95% Conf Intervals Lower	idence Upper
Death	6.9%	1.2%	5.6*	1.61	19.34
CHF Hospitalization (during DB)	5.3%	2.0%	2.7	0.89	8.39

b: during double-blind therapy or was related to Serious AE during double-blind

*: odds are greater for placebo group as compared to losartan group

TABLE 6—Meta-Analysis[@] CLINICAL EVENTS—PROTOCOLS A &B

	PBO (N=245)	Losartan (N=491)	Odds Ratio	95% Conf Intervals Lower	idence Upper
Death	5.3%	1.4%	3.5*	1.43	8.77
CHF Hospitalization (during double-blind)	6.9%	3.1% ª	2.4*	1.19	4.76

- @: Peto-modified Mantel-Haenszel method
- 5 b: during double-blind therapy or was related to Serious AE during double-blind
 - a: one patient discontinued due to worsening HF and was hospitalized 2 days after discontinuation from double-blind.
 - *: odds are greater for placebo group as compared to losartan group

WHAT IS CLAIMED IS:

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1. A method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist.

- 2. A method for reducing mortality by administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458, SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498, andYM31472.
- 3. The method as recited in Claim 2, wherein the angiotensin II receptor antagonist is selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

A method for reducing mortality by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

$$\begin{array}{c|c}
 & R^7 \\
 & R^6 \\
 & R^8 \\
 & (CH_2)_r \\
 & R^1 \\
 & R^2 \\
 & R^3
\end{array}$$

5 wherein:

 $\label{eq:conhomo} \begin{array}{c} \mathsf{CO_2H} \\ \text{4-CONHNHSO}_2\mathsf{CF}_3; \text{4-CONH-CHCH}_2\mathsf{C}_6\mathsf{H}_5 \end{array}; \\ \text{$(F$ isomer)}$

$$\begin{array}{c} O \\ -C-NHSO_2 - (CH_2)_s \end{array} ;$$

R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃;

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl - 76 -

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or cycloalkylalkynyl 5 to 10 carbon atoms; $(CH_2)_s Z(CH_2)_m R^5$ optionally substituted with F or CO_2R^{14} ; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

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R7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where v=1-6, C₆F₅; CN;

The straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

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R⁸ is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH₂)_m-imidazol-1-yl; -(CH₂)_m-1,2,3-triazolyl optionally substituted with one or two group selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms; -(CH₂)_s tetrazolyl;

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-(CH₂)_{n-1}CH-R¹¹; -(CH₂)_nOCR¹⁴; -(CH₂)_nSR¹⁵; OR¹⁷

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R14 O O -CH=CH(CH2)sCHOR15; -CH=CH(CH2)sCR16; -CR16;

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O -CH=CH(CH₂)_sOCR¹¹; (CH₂)_s-CH-COR¹⁶; CH₃

O Y -(CH₂)_nCR₁₆; -(CH₂)_nOCNHR₁₀; -(CH₂)_nNR₁₁COR₁₀;

5 -(CH₂) $_m$ NO₂; -CH=N-NR¹¹R¹⁷;

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow (CH_{2})_{n}-N \longrightarrow (CH_{3}O) ;$$

$$-(CH_{2})_{n-1}C-N \longrightarrow (CH_{3}O) ;$$

$$-(CH_{3}O) \longrightarrow (CH_$$

R24 O R9 is: -CH-OCR21;

10 R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)pC6H5;

R11 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

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R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

-PO3H2; -C(CF3)2OH; -NHSO2CH3; -NHSO2CF3;

-NHCOCF3; -CONHOR12; -SO2NH2;

- R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
- R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;
- R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
 - R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 $Q;$

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Q is NR²⁰, O or CH₂;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or - CHCH2CO2CH3;
NH2

R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

10 R²⁴ is H, CH₃ or -C₆H₅;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

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- R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
- R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;

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- R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH₂)q-;
- R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;

X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23

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-NHC(R²⁷)(R²⁸)-, -NR²³SO₂-, -SO₂NR²³-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH₂CH₂-, -C(R²⁷)(R²⁸)NH-,

Y is O or S;

Z is O, NR11, or S;

5 m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

q is 2 to 3;

r is 0 to 2;

10 s is 0 to 5;

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t is 0 or 1;

and pharmaceutically acceptable salts of these compounds; provided that:

(1) the R¹ group is not in the ortho position;

(2) when R^1 is

$$-x - \begin{cases} R^{13} \\ - R^{2} \end{cases}$$

X is a single bond, and R13 is CO₂H, or

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

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(3) when R^1 is

$$-x - \begin{cases} R^{13} \\ -R^{3} \end{cases}$$

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and X is other than a single bond, then R^{13} must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

- (4) when R¹ is 4-CO₂H or a salt thereof, R⁶ cannot be S-alkyl;
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- (5) when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
- (6) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ \\ \\ R^2 \end{array} \right)$$

X is -OCH2-, and R^{13} is 2-CO2H, and R^7 is H then R^6 is not C2H5S;

(7) when R^1 is

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and R6 is n-hexyl then R7 and R8 are not both hydrogen;

10 (8) when R^1 is

R6 is not methoxybenzyl;

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- (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
- (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

X is -NH-C-, R13 is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^3 \end{array} \right)$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

$$R^1 = X - (R^{13}) + (R^{13}) +$$

X is a single bond, R7 is Cl, and R8 is -CHO, then R13 is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

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X is a single bond, R7 is C1, and R8 is -CHO, then R13 is not 4-(tetrazol-5-yl).

5. The method as recited in Claim 4, using the angiotensin II receptor antagonist of formula I:

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wherein:

R1 is -CO2H; -NHSO2CF3;

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R6 is alkyl of 3 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, alkynyl of 3 to 10 carbon atoms, cycloalkyl of 3 to 8 carbon atoms, benzyl substituted on the phenyl ring with up to two groups selected from alkoxy of 1 to 4 carbon atoms, halogen, alkyl of 1 to 4 carbon atoms, and nitro;

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R8 is phenylalkenyl wherein the aliphatic portion is 2 to 4 carbon atoms, $-(CH_2)_m$ -imidazol-1yl, $-(CH_2)_m$ 1,2,3-triazolyl optionally substituted with one or two groups selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms,

(CH₂)_m-tetrazolyl, -(CH₂)_nOR11; -(CH₂)_nOCR14; R14 -CH=CH(CH₂)_sCR16, -CH=CH(CH₂)_sCHOR15; O O O -(CH₂)_nCR16; -(CH₂)_nNHCOR10; -(CH₂)_nNHSO₂R10; O (CH₂)_mF; -CR16; 10 R13 is -CO₂H, -CO₂R⁹, NHSO₂CF3; SO₃H; N=N N ;

R16 is H, alkyl of 1 to 5 carbon atoms, OR17, or NR18R19;

X is carbon-carbon single bond, -CO-, -CON-, -CH2CH2-, -NCO-, R23 R23

-OCH2-, -CH2O-, -SCH2-, -CH2S-, -NHCH2-, -CH2NH- or -CH=CH-; and pharmaceutically acceptable salts of these compounds.

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6. The method as recited in Claim 5, using the angiotensin II receptor antagonist of formula I wherein:

R2 is H, alkyl of 1 to 4 carbon atoms, halogen, or alkoxy of 1 to 4 carbon atoms;

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R6 is alkyl, alkenyl or alkynyl of 3 to 7 carbon atoms;

R7 is H, Cl, Br, $C_{\nu}F_{2\nu+1}$, where v=1-3, or -CR16;

5 R10 is CF3, alkyl of 1 to 6 carbon atoms or phenyl;

R11 is H, or alkyl of 1 to 4 carbon atoms;

R13 is CO2H; CO2CH2OCOC(CH3)3; NHSO2CF3;

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R14 is H, or alkyl of 1 to 4 carbon atoms;

R15 is H, alkyl of 1 to 4 carbon atoms, or acyl of 1 to 4 carbon atoms;

R16 is H, alkyl of 1 to 5 carbon atoms; OR17; or

$$-N$$

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m is 1 to 5;

X is single bond, -O-; -CO-; -NHCO-; or -OCH2-; and pharmaceutically acceptable salts.

- 7. The method as recited in Claim 4, wherein the angiotensin II receptor antagonist of formula I is selected from the group consisting of:
 - 2-Butyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-

(hydroxymethyl)imidazole.

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- 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-[(methoxy-carbonyl)aminomethyl]imidazole.
- 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-[(propoxy-carbonyl)aminomethyl]imidazole.
- 2-Butyl-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]imidazole-5-carboxaldehyde.
- 2-Butyl-1-[(2'-carboxybiphenyl-4-yl)methyl]-imidazole-5-carbox-aldehyde.
 - 2-(1E-Butenyl)-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
 - 2-(1E-Butenyl)-4-chloro-1-[(2'-carboxybiphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.
 - 2-Propyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
 - 2-Propyl-4-chloro-1[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.
- 20 2-Butyl-4-chloro-1-[2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-imidzole-5-carboxaldehyde.
 - 2-(1E-Butenyl)-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-hydroxymethyl)imidazole.
 - 2(1E-Butenyl)-4-chloro-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxaldehyde.
 - 2-Butyl-4-chloro-1-[(2'-(1H-tetrazol-5-yl)-biphenyl-4-yl)methyl]-imidazole-5-carboxylic acid.
 - 2-Propyl-4-chloro-1-[(2-'(1H-tetrazol-5-yl)-biphenyl-4-yl)methyl]-imidazole-5-carboxylic acid.
- 30 2-Propyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.
 - 2-Propyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxylmethyl)imidazole.
 - 2-Butyl-4-trifluoromethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.

2-Propyl-4-trifluoromethyl-1-[(2'-(carboxybiphenyl-4-yl)methyl]-imidazole-5-carboxaldehyde.

- 2-Propyl-4-pentafluoroethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]-5-(hydroxymethyl)imidazole.
- 5 2-Propyl-1-[(2-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-4,5,-dicarboxylic acid.
 - 2-Propyl-4-pentafluoroethyl-1-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxylic acid.
- 2-Propyl-4-pentafluoroethyl-[(2'-(1H-tetrazol-5-yl)biphenyl-4-yl)methyl]imidazole-5-carboxaldehyde, or a pharmaceutically acceptable salt thereof.
 - 8. The method as recited in Claim 7, wherein the angiotensin II receptor antagonist of formula I is:
- 15 2-Butyl-4-chloro-1-[(2'-tetrazol-5-yl)biphenyl-4-yl]methyl]-5-(hydroxy-methyl)imidazole; and
 - 2-Butyl-4-chloro-1-[(2'-tetrazol-5-yl)biphenyl-4-yl]methylimidazole-5-carboxylic acid or a pharmaceutically acceptable salt thereof.
- 9. The method as recited in Claim 8, wherein the symptomatic heart failure patient is 65 years of age or older.
- 10. A method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist.
- administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458,
- 35 SL910102, UP2696, YM358, EMD66397, ME3221, TAK536,

BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498, andYM31472.

- 12. The method as recited in Claim 11, wherein the angiotensin II receptor antagonist is selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.
- 13. A method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

wherein:

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R¹ is:

R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃;

CONHOR¹²;
$$SO_2NH_2$$
; $N-N$ N; aryl; or furyl; N

5

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

10

- R⁵ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;
- R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO2R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl or cycloalkylalkynyl 5 to 10 carbon atoms; (CH2)_SZ(CH2)_mR⁵ optionally substituted with F or CO2R¹⁴; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

R⁷ is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where v=1-6, C6F5; CN;

25

O

C-R¹⁶; straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH₃, CF₃, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

	R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10
	carbon atoms, or the same groups substituted with F;
	phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; $-(CH_2)_m$ -imidazol-1-yl; $-(CH_2)_m$ -1,2,3-triazolyl
5.	optionally substituted with one or two group selected from CO ₂ CH ₃ or alkyl of 1 to 4 carbon atoms; -(CH ₂) _s
	tetrazolyl;
	Q
	-(CH ₂) _{n-1} CH-R ¹¹ ; -(CH ₂) _n OCR ¹⁴ ; -(CH ₂) _n SR ¹⁵ ;
10	OR17
	\mathbb{R}^{14} Q Q
	R ¹⁴ O O -CH=CH(CH ₂) _s CHOR ¹⁵ ; -CH=CH(CH ₂) _s CR ¹⁶ ; -CR ¹⁶ ;
•	Q
. •	-CH=CH(CH ₂) _s OCR ¹¹ ; (CH ₂) _s -CH-COR ¹⁶ ;
15	ĆH3
	O Y -(CH ₂) _n CR ¹⁶ ; -(CH ₂) _n OCNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ COR ¹⁰ ;
	O -(CH ₂) _n NR ¹¹ CNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ SO ₂ R ¹⁰ ;
20	Y -(CH ₂) _n NR ¹¹ CR ¹⁰ ; -(CH ₂) _m F; -(CH ₂) _m ONO ₂ ; -CH ₂ N ₃ ; -(CH ₂) _m NO ₂ ; -CH=N-NR ¹¹ R ¹⁷ ;
	-(Cn2)mi(O2, -Cn=14-14K-1K-1,

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow NH ;$$

$$-(CH_{2})_$$

- R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH₂)_pC6H₅;
- R¹¹ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

5

R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;

10

R16 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH2)_pC6H5, OR17, or NR18R19;

R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

15

R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 $(CH2)$

- Q is NR20, O or CH2;
- R20 is H, alkyl of 1-4 carbon atoms, or phenyl;
- 25 R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or CHCH2CO2CH3;
 NH2

R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

5 R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

- 10 R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
 - R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;
- R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;
 - R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;

20
X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-,

R26 R23 R23

-NHC(R²⁷)(R²⁸)-, -NR²³SO₂-, -SO₂NR²³-, -CH=CH-, -

CF=CF-, -CH=CF-, -CF=CH-, -CH2CH2-, -C(R²⁷)(R²⁸)NH-,

Y is O or S;

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Z is O, NR11, or S;

m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

5 q is 2 to 3;

r is 0 to 2;

s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds;

10 provided that:

- (1) the R^1 group is not in the ortho position;
- (2) when R1 is

$$-X - \begin{bmatrix} R^{13} \\ -R^{2} \end{bmatrix}$$

15

X is a single bond, and R13 is CO2H, or

20

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

(3) when R1 is

$$-X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

5 (4) when R1 is 4-CO₂H or a salt thereof, R6 cannot be S-alkyl;

(5) when R1 is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;

(6) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ \\ \\ R^2 \end{array} \right)$$

X is -OCH₂-, and R¹³ is 2-CO₂H, and R⁷ is H then R⁶ is not C₂H₅S;

(7) when R^1 is

CF₃SO₂HN
—CONH—

20

and R6 is n-hexyl then R7 and R8 are not both hydrogen;

(8) when R1 is

5

R6 is not methoxybenzyl;

- 10 (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
 - (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^2 \end{array} \right)$$

Q X is -NH-C-, R¹³ is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

O X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO₂CH₃;

(12) when r=1,

$$R^1 = X - (R^{13})$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \begin{cases} R^{13} \\ R^2 \end{cases}$$

20

10

15

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 4-(tetrazol-5-yl).

14. The method as recited in Claim 13, wherein the angiotensin II receptor antagonist is losartan or EXP-3174, or a pharmaceutically acceptable salt thereof.

- 5 15. The method as recited in Claim 14, wherein the symptomatic heart failure patient is 65 years of age or older.
- 16. A method for reducing mortality and sudden cardiac death by administering to a symptomatic heart failure patients a
 10 therapeutically effective amount of an angiotensin II receptor antagonist.
- 17. A method for reducing mortality and sudden cardiac death by administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458,
- SL910102, UP2696, YM358, EMD66397, ME3221, TAK536,
 BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684,
 EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057,
 LY266099, LY301875, PD123177, PD126055, SC51757, SC54629,
 U96849, UK77778, WAY126227, WK1260, WK1492, YH1498,
 andYM31472.
- 18. The method as recited in Claim 16, wherein the angiotensin II receptor antagonist is selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.
 - 19. A method for reducing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II antagonist formula I

$$R^6$$
 N
 R^7
 R^8
 $(CH_2)_r$
 R^1
 R^2
 R^3

wherein: R¹ is:

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R² is H; Cl; Br; I; F; NO₂; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO₂H; CO₂R⁹; HNSO₂CH₃; NHSO₂CF₃; CONHOR¹²; SO₂NH₂; N-N, aryl; or furyl;

5

R³ is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO₂ or CO₂R11;

10

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or 15 CO2R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl or cycloalkylalkynyl 5 to 10 carbon atoms; (CH2)_sZ(CH2)_mR⁵ optionally substituted with F or CO₂R¹⁴; benzyl substituted on the phenyl ring with 1 or 2 20 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4

carbon atoms or nitro;

 R^7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where $\nu=1-6$, C₆F₅; CN;

25

—Ü-R¹⁶; straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

	R ⁸ is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH ₂) _m -imidazol-1-yl; -(CH ₂) _m -1,2,3-triazolyl
5	optionally substituted with one or two group selected from CO ₂ CH ₃ or alkyl of 1 to 4 carbon atoms; -(CH ₂) _s
	tetrazolyl;
10	O -(CH ₂) _{n-1} CH-R ¹¹ ; -(CH ₂) _n OCR ¹⁴ ; -(CH ₂) _n SR ¹⁵ ; OR ¹⁷
	R ¹⁴ O O -CH=CH(CH ₂), CHOR ¹⁵ ; -CH=CH(CH ₂), CR ¹⁶ ; -CR ¹⁶ ;
15	O -CH=CH(CH ₂) ₅ OCR ¹¹ ; (CH ₂) ₅ -CH-COR ¹⁶ ; CH ₃
	O Y -(CH ₂) _n CR ¹⁶ ; -(CH ₂) _n OCNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ COR ¹⁰ ;
	O -(CH ₂) _n NR ¹¹ CNHR ¹⁰ ; -(CH ₂) _n NR ¹¹ SO ₂ R ¹⁰ ;
20	Y -(CH ₂) _n NR ¹¹ CR ¹⁰ ; -(CH ₂) _m F; -(CH ₂) _m ONO ₂ ; -CH ₂ N ₃ ; -(CH ₂) _m NO ₂ ; -CH=N-NR ¹¹ R ¹⁷ ;

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow (CH_{2})_{s} \longrightarrow (CH_{2})_{s} \longrightarrow (CH_{2})_{n}-N \longrightarrow (CH_{2})_{n}-N \longrightarrow (CH_{3})_{n} \longrightarrow (CH_{3})_{n}-N \longrightarrow (CH_{3})_{$$

- 5 R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)_pC6H5;
- R11 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

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5

15

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} N-N \\ N\end{array} \end{array} \\ \begin{array}{c} N-N \\ N\end{array} \end{array} \\ \begin{array}{c} -CH_2 \\ N \\ N\end{array} \end{array} \begin{array}{c} N-N \\ N \end{array} \\ \begin{array}{c} -CONH \\ N \\ N\end{array} \end{array} \begin{array}{c} N-N \\ N \end{array} \\ \begin{array}{c} Or \\ N=N \end{array} \end{array} \begin{array}{c} N=N \\ N=N \end{array}$$

- R¹⁴ is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- R¹⁵ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
- 10 R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;
 - R¹⁷ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
 - R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

20 Q is NR20, O or CH2;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

25 R²¹ is alkyl of 1 to 6 carbon atoms, -NR²²R²³, or - CHCH₂CO₂CH₃; NH₂

 R^{22} and R^{23} independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

5 R²⁴ is H, CH₃ or -C₆H₅;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

- 10 R²⁶ is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl:
 - R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;
- 15 R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;
 - R³¹ is H, alkyl or 1 to 4 carbon atoms, -CH₂CH=CH₂ or -CH₂C6H₄R³²;

20
X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-,

-NHC(R27)(R28)-, -NR23SO2-, -SO2NR23-, -CH=CH-, -

$$CF = CF -, -CH = CF -, -CF = CH -, -CH_2CH_2 -, -C(R^{27})(R^{28})NH -,$$

$$-CF_2CF_2$$
; OR^{14} $OCOR^{17}$ NR^{25} ; $-CH^-$;

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Y is O or S;

Z is O, NR11, or S;

m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

5 q is 2 to 3;

r is 0 to 2;

s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds;

10 provided that:

- (1) the R¹ group is not in the ortho position;
- (2) when R¹ is

$$-x - \begin{cases} R^{13} \\ -R^{3} \end{cases}$$

15

X is a single bond, and R13 is CO₂H, or

20

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

(3) when R^1 is

$$-X - \left(\frac{R^{13}}{R^2} \right)^{13}$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

5

10

- (4) when R¹ is 4-CO₂H or a salt thereof, R⁶ cannot be S-alkyl;
- (5) when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
 - (6) when R1 is

$$-x = \begin{bmatrix} R^{13} \\ R^3 \end{bmatrix}$$

15

X is -OCH2-, and R^{13} is 2-CO2H, and R^7 is H then R^6 is not C2H5S;

(7) when R^1 is

$$\begin{array}{c} \text{CF}_3\text{SO}_2\text{HN} \\ -\text{CONH} - \end{array},$$

and R6 is n-hexyl then R7 and R8 are not both hydrogen;

(8) when R^1 is

5

R6 is not methoxybenzyl;

- 10 (9) the R⁶ group is not -CHCH2CH3 or CH2OH;
 - (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^3 \end{array} \right)$$

X is -NH-C-, R13 is 2-NHSO2CF3, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right),$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

$$R^1 = X - (R^{13})$$

X is a single bond, R7 is Cl, and R8 is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - (R^{13})$$

20

10

X is a single bond, R^7 is Cl, and R^8 is -CHO, then R^{13} is not 4-(tetrazol-5-yl).

20. The method as recited in Claim 19, wherein the angiotensin II receptor antagonist is losartan or EXP-3174 or its pharmaceutically acceptable salt thereof.

- 5 21. The method as recited in Claim 20, wherein the symptomatic heart failure patient is 65 years of age or older.
- 22. A method for preventing sudden cardiac death by administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist.
- A method for preventing sudden cardiac death by 23. administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, 15 losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1.1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3Himidazo[4.5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458, 20 SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498,
 - 24. The method as recited in Claim 23, wherein the angiotensin II receptor antagonist is selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.

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andYM31472.

25. A method for preventing sudden cardiac death by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

PCT/US98/00534 WO 98/30216

$$R^{6}$$
 N
 R^{8}
 $(CH_{2})_{r}$
 R^{1}
 R^{2}
 R^{3}

wherein:

$$CO_2H$$

4-CONHNHSO $_2CF_3$; 4-CONH $-CHCH_2C_6H_5$; (*F*isomer)

$$HO_2C$$
 R^{11}
 $A-CON$
 HO_2C
 R^{11}
 HO_2C
 HO

$$A = N$$
 $A = N$
 $A =$

$$\begin{array}{c} {\rm O} \\ {\rm -C-NHSO_2---} \ ({\rm CH_2})_{\rm s} \end{array} ;$$

R2 is H; Cl; Br; I; F; NO2; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO2H; CO2R9; HNSO2CH3; NHSO2CF3;

R3 is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R⁴ is CN, NO₂ or CO₂R¹¹;

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl - 115 -

5

or cycloalkylalkynyl 5 to 10 carbon atoms; $(CH_2)_s Z(CH_2)_m R^5$ optionally substituted with F or $CO_2R_1^{14}$; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

5

R7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where $\nu=1-6$, C6F5; CN;

10

straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

15

R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH₂)_m-imidazol-1-yl; -(CH₂)_m-1,2,3-triazolyl optionally substituted with one or two group selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms; -(CH₂)_s tetrazolyl;

20

-(CH₂)_{n-1}CH-R¹¹; -(CH₂)_nOCR¹⁴; -(CH₂)_nSR¹⁵; OR¹⁷

25

R14 O O O -CH=CH(CH2)sCHOR15; -CH=CH(CH2)sCR16; -CR16;

30

-CH=CH(CH₂)_sOCR¹¹; (CH₂)_s-CH-COR¹⁶; CH₃

O Y Y Y - (CH₂)_nCR₁₆; -(CH₂)_nOCNHR₁₀; -(CH₂)_nNR₁₁COR₁₀;

5

 $O_{-(CH_2)_nNR_{11}CNHR_{10}; -(CH_2)_nNR_{11}SO_2R_{10};}$

-(CH₂)_nNR₁₁CR₁₀; -(CH₂)_mF; -(CH₂)_mONO₂; -CH₂N₃; -(CH₂)_mNO₂; -CH=N-NR₁₁R₁₇;

$$-(CH_2)_s$$
 N
 CF_3 ; $-(CH_2)_n$
 N
 CH_3O

$$\longrightarrow (CH_2)_{n-1}C - N \longrightarrow N \longrightarrow ;$$

$$CH_3O$$

$$-CH=N-NH-SO_2-$$
 ; or $-CH=N-NH-$

R²⁴ O R⁹ is: -CH-OCR²¹

- 10 R¹⁰ is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH₂)_pC₆H₅;
- R¹¹ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

R13 is -CO₂H; -CO₂R⁹; -CH₂CO₂H, -CH₂CO₂R⁹;

-PO₃H₂; -C(CF₃)₂OH; -NHSO₂CH₃; -NHSO₂CF₃; -NHCOCF₃; -CONHOR¹²; -SO₂NH₂;

R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R¹⁵ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;

R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;

R¹⁷ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R¹⁸ and R¹⁹ independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

Q is NR²⁰, O or CH₂;

5

10

15

20

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or - CHCH2CO2CH3; NH2

 R^{22} and R^{23} independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as $(CH_2)_u$, where u is 3-6;

10 R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

15

5

- R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
- R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;

- R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;
- R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;
- X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23

 -NHC(R²⁷)(R²⁸)-, -NR²³SO₂-, -SO₂NR²³-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH₂CH₂-, -C(R²⁷)(R²⁸)NH-,

Y is O or S;

Z is O, NR11, or S;

5 m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

q is 2 to 3;

r is 0 to 2;

10 s is 0 to 5;

15

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds; provided that:

- (1) the R¹ group is not in the ortho position;
- (2) when R1 is

$$-x - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

20 X is a single bond, and R13 is CO₂H, or

then R¹³ must be in the ortho or meta position; or when R¹ and X are as above and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

5

(3) when R¹ is

$$-x - \begin{cases} R^{13} \\ R^{2} \end{cases}$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

10

(4) when R1 is 4-CO₂H or a salt thereof, R6 cannot be S-alkyl;

15

- (5) when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
- (6) when R1 is

$$-X - \left(\begin{array}{c} R^{13} \\ \\ \\ R^2 \end{array} \right)$$

X is -OCH₂-, and R^{13} is 2-CO₂H, and R^7 is H then R^6 is not C₂H₅S;

(7) when R^1 is

5

and R6 is n-hexyl then R7 and R8 are not both hydrogen;

10 (8) when R^1 is

R6 is not methoxybenzyl;

15

- (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
- (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

Q X is -NH-C-, R¹³ is 2-NHSO₂CF₃, and R⁶ is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ = \end{array} \right)$$

$$R^{2}$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO2CH3;

(12) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

20

15

10

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 4-(tetrazol-5-yl).

26. The method as recited in Claim 25, wherein the angiotensin II receptor antagonist is losartan or EXP-3174, or a pharmaceutically acceptable thereof.

- 5 27. The method as recited in Claim 26, wherein the symptomatic heart failure patient is 65 years of age or older.
- 28. A method for reducing hospitalization by administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist.
- 29. A method for reducing hospitalization by administering to a symptomatic heart failure patients a therapeutically effective amount of an angiotensin II receptor antagonist selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698, 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine, BAY106734, BIBR363, CL329167, E4177, EMD73495, HN65021, HR720, HOE720, LRB081, SC52458,
- 20 SL910102, UP2696, YM358, EMD66397, ME3221, TAK536, BMS184698, CGP42112A, CGP49870, CP148130, E4188, EMD66684, EXP9954, FR1153332, GA0050, KT3579, LF70156, LRB057, LY266099, LY301875, PD123177, PD126055, SC51757, SC54629, U96849, UK77778, WAY126227, WK1260, WK1492, YH1498,
- 25 and YM 31472.
- 30. The method as recited in Claim 29, wherein the angiotensin II receptor antagonist is selected from the group consisting of: candesartan cilexetil, eprosartan, irbesartan, losartan, tasosartan, telmisartan, valsartan, BMS-184698 and 3-(2'-(tetrazol-5-yl)-1,1'-biphen-4-yl)methyl-5,7-dimethyl-2-ethyl-3H-imidazo[4,5-b]pyridine.
- 31. A method for reducing hospitalization by administering to a symptomatic heart failure patient a therapeutically effective amount of an angiotensin II receptor antagonist of formula I

wherein:

$$CO_2H$$

4-CONHNHSO $_2CF_3$; 4-CONH $-CHCH_2C_6H_5$; (*F*isomer)

$$HO_2C$$
 R^{11}
 HO_2C
 $HO_$

$$\begin{array}{c} O \\ -\text{C-NHSO}_2 - - (\text{CH}_2)_s \end{array} ;$$

R2 is H; Cl; Br; I; F; NO2; CN; alkyl of 1 to 4 carbon atoms; acyloxy of 1 to 4 carbon atoms; alkoxy of 1 to 4 carbon atoms; CO2H; CO2R9; HNSO2CH3; NHSO2CF3;

R3 is H; Cl, Br, I or F; alkyl of 1 to 4 carbon atoms or alkoxy of 1 to 4 carbon atoms;

R4 is CN, NO2 or CO2R11;

R5 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms alkenyl or alkynyl of 2 to 4 carbon atoms;

R6 is alkyl of 2 to 10 carbon atoms, alkenyl or alkynyl of 3 to 10 carbon atoms or the same groups substituted with F or CO₂R¹⁴; cycloalkyl of 3 to 8 carbon atoms, cycloalkylalkyl, of 4 to 10 carbon atoms; cycloalkylalkenyl - 126 -

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5

10

or cycloalkylalkynyl 5 to 10 carbon atoms; $(CH_2)_s Z(CH_2)_m R^5$ optionally substituted with F or CO_2R^{14} ; benzyl substituted on the phenyl ring with 1 or 2 halogens, alkoxy of 1 to 4 carbon atoms, alkyl of 1 to 4 carbon atoms or nitro;

5

 R^7 is H, F, Cl, Br, I, NO₂, $C_{\nu}F_{2\nu+1}$, where $\nu=1-6$, C6F5; CN;

The straight or branched alkyl of 1 to 6 carbon atoms; phenyl or phenylalkyl, where alkyl is 1 to 3 carbon atoms; or substituted phenyl or substituted phenylalkyl, where alkyl is 1 to 3 carbon atoms, substituted with one or two substituents selected from alkyl of 1 to 4 carbon atoms, F, Cl, Br, OH, OCH3, CF3, and COOR, where R is H, alkyl of 1 to 4 carbon atoms, or phenyl;

15

10

R8 is H, CN, alkyl of 1 to 10 carbon atoms, alkenyl of 3 to 10 carbon atoms, or the same groups substituted with F; phenylalkenyl wherein the aliphatic portion is 2 to 6 carbon atoms; -(CH₂)_m-imidazol-1-yl; -(CH₂)_m-1,2,3-triazolyl optionally substituted with one or two group selected from CO₂CH₃ or alkyl of 1 to 4 carbon atoms; -(CH₂)_s tetrazolyl;

20

-(CH₂)_{n-1}CH-R¹¹; -(CH₂)_nOCR¹⁴; -(CH₂)_nSR¹⁵; O_{R}^{17}

25

R14 O O -CH=CH(CH₂)sCHOR¹⁵; -CH=CH(CH₂)sCR¹⁶; -CR¹⁶;

30

-CH=CH(CH₂)_sOCR¹¹; (CH₂)_s-CH-COR¹⁶; CH₃

O Y Y Y -(CH₂)_nCR¹⁶; -(CH₂)_nOCNHR¹⁰; -(CH₂)_nNR¹¹COR¹⁰;

5

O -(CH₂)_nNR₁₁C_NHR₁₀; -(CH₂)_nNR₁₁SO₂R₁₀; Y -(CH₂)_nNR₁₁C_R10; -(CH₂)_mF; -(CH₂)_mONO₂; -CH₂N₃; -(CH₂)_mNO₂; -CH=N-NR₁₁R₁₇;

$$-(CH_{2})_{m}-N \longrightarrow (CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{s} \longrightarrow NH ;$$

$$-(CH_{2})_{n}-N \longrightarrow NH ;$$

$$-(CH_{2})_$$

R24 O R9 is: -CH-OCR21;

10
R10 is alkyl of 1 to 6 carbon atoms or perfluoroalkyl of 1 to 6 carbon atoms, 1-adamantyl, 1-naphthyl, 1-(1-naphthyl)ethyl, or (CH2)pC6H5;

15 R11 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;

R12 is H, methyl or benzyl;

5

- 10 R14 is H, alkyl or perfluoroalkyl of 1 to 8 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- R15 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl, benzyl, acyl of 1 to 4 carbon atoms, phenacyl;
 - R¹⁶ is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, (CH₂)_pC₆H₅, OR¹⁷, or NR¹⁸R¹⁹;
- 20 R17 is H, alkyl of 1 to 6 carbon atoms, cycloalkyl of 3 to 6 carbon atoms, phenyl or benzyl;
- R18 and R19 independently are H, alkyl of 1 to 4 carbon atoms, phenyl, benzyl, α-methylbenzyl, or taken together with the nitrogen form a ring of the formula

$$-N$$
 Q ;

Q is NR20, O or CH2;

R20 is H, alkyl of 1-4 carbon atoms, or phenyl;

- 5 R21 is alkyl of 1 to 6 carbon atoms, -NR22R23, or CHCH2CO2CH3; NH2
- R22 and R23 independently are H, alkyl of 1 to 6 carbon atoms, benzyl, or are taken together as (CH₂)_u, where u is 3-6;

R24 is H, CH3 or -C6H5;

R25 is NR27R28, OR28, NHCONH2, NHCSNH2,

—
$$NHSO_2$$
—C H_3 ; or — $NHSO_2$ —

15

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- R26 is hydrogen, alkyl with from 1 to 6 carbon atoms, benzyl, or allyl;
- R27 and R28 are independently hydrogen, alkyl with from 1 to 5 carbon atoms, or phenyl;
- R29 and R30 are independently alkyl of 1-4 carbon atoms or taken together are -(CH2)q-;
- 25 R31 is H, alkyl or 1 to 4 carbon atoms, -CH2CH=CH2 or -CH2C6H4R32;
- X is a carbon-carbon single bond, -CO-, -CH₂-, -O-, -S-, -NH-, -N-, -CON-, -NCO-, -OCH₂-, -CH₂O-, -SCH₂-, -CH₂S-, R26 R23 R23 -NHC(R²7)(R²8)-, -NR²3SO₂-, -SO₂NR²3-, -CH=CH-, -CF=CF-, -CH=CF-, -CF=CH-, -CH₂CH₂-, -C(R²7)(R²8)NH-,

$$-CF_2CF_2$$
; OR^{14} $OCOR^{17}$ NR^{25} $-CH^{-}$; $-CH^{-}$; $-CH^{-}$; $-CH^{-}$; or $-CH^{-}$;

Y is O or S;

Z is O, NR11, or S;

5 m is 1 to 5;

n is 1 to 10;

p is 0 to 3;

q is 2 to 3;

r is 0 to 2;

10 s is 0 to 5;

t is 0 or 1;

and pharmaceutically acceptable salts of these compounds; provided that:

(1) the R¹ group is not in the ortho position;

15

20

(2) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ - \\ R^{2} \end{array} \right)$$

X is a single bond, and R13 is CO₂H, or

then R13 must be in the ortho or meta position; or when R1 and X are as above and R13 is NHSO₂CF₃ or NHSO₂CH₃, R¹³ must be ortho;

(3) when R^1 is

$$-X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

and X is other than a single bond, then R¹³ must be ortho except when X=NR²³CO and R¹³ is NHSO₂CF₃ or NHSO₂CH₃, then R¹³ must be ortho or meta;

- (4) when R1 is 4-CO2H or a salt thereof, R6 cannot be S-alkyl;
- when R¹ is 4-CO₂H or a salt thereof, the substituent on the 4-position of the imidazole cannot be CH₂OH, CH₂OCOCH₃, or CH₂CO₂H;
 - (6) when R¹ is

$$-X - \left(\begin{array}{c} R^{13} \\ R^{2} \end{array} \right)$$

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X is -OCH₂-, and R^{13} is 2-CO₂H, and R^{7} is H then R^{6} is not C₂H₅S;

(7) when R^1 is

5

and R6 is n-hexyl then R7 and R8 are not both hydrogen;

10 (8) when R^1 is

R6 is not methoxybenzyl;

15

- (9) the R6 group is not -CHCH2CH2CH3 or CH2OH;
- (10) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ - \\ R^2 \end{array} \right)$$

X is -NH-C-, R13 is 2-NHSO₂CF₃, and R6 is n-propyl, then R⁷ and R⁸ are not -CO₂CH₃;

5 (11) when r=0, R^1 is

$$X - \left(\begin{array}{c} R^{13} \\ R^3 \end{array} \right)$$

X is NH-C-, R13 is 2-COOH, and R6 is n-propyl, then R7 and R8 are not -CO₂CH₃;

(12) when r=1,

$$R^1 = X - (R^{13}) + (R^{13}) +$$

X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 3-(tetrazol-5-yl);

(13) when r=1,

$$R^1 = X - \left(\begin{array}{c} R^{13} \\ + \\ R^2 \end{array} \right)$$

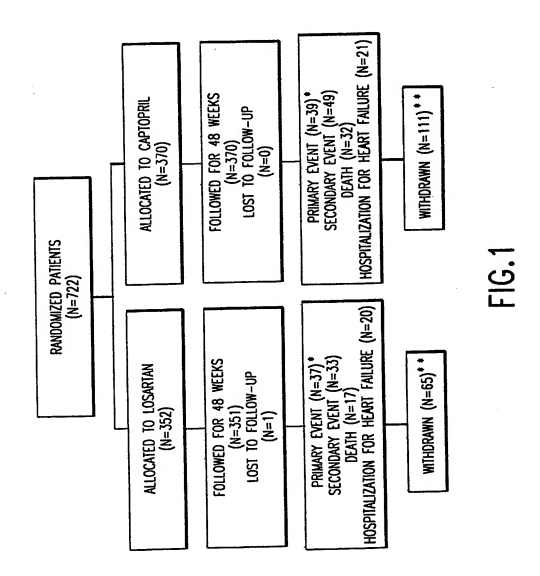
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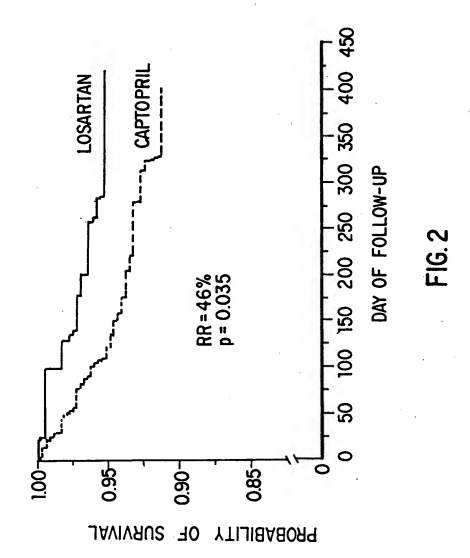
X is a single bond, R⁷ is Cl, and R⁸ is -CHO, then R¹³ is not 4-(tetrazol-5-yl).

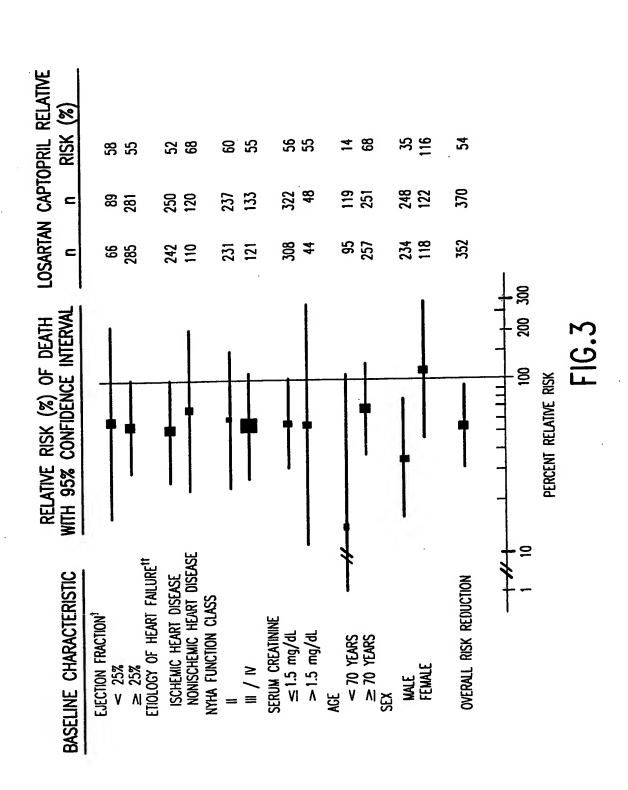
32. The method as recited in Claim 31, wherein the angiotensin II receptor antagonist is losartan or EXP-3174, or its pharmaceutically acceptable salt thereof.

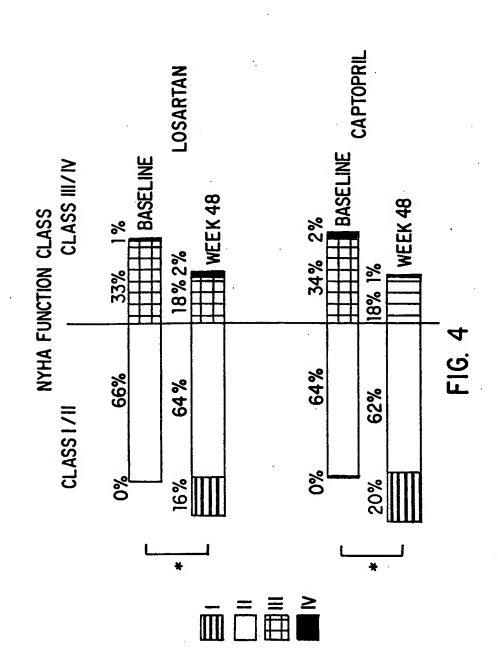
5 33. The method as recited in Claim 32, wherein the symptomatic heart failure patient is 65 years of age or older.

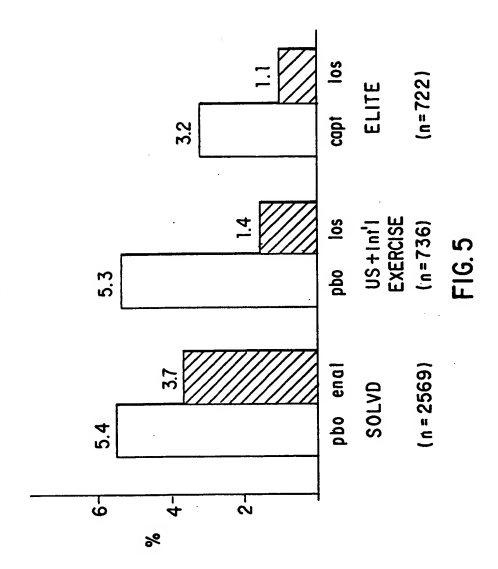


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יייכיסטורי יייט טמטטיביי

Intern. unal Application No PCT/US 98/00534

A. CLASSIFICATION OF SUBJECT MATTER
1PC 6 A61K31/41 A61K31/415 A61K31/44 A61K31/47 A61K31/505
A61K38/04

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 6 A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUM	NTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X .	O. LENZ ET AL.: "Effects of angiotensin II and angiotensin-converting enzyme inhibitors on human myocardium." EUR. J. PHARMACOL., vol. 294, no. 1, 1995, pages 17-27, XP002065907 see the whole document	1-8, 10-14, 16-20, 22-26, 28-32
X	N.A. AWAN ET AL.: "Direct selective blockade of the vascular angiotensin II receptors in therapy for hypertension and severe congestive heart failure." AM. HEART JOURNAL, vol. 131, no. 1, 1996, pages 177-185, XP002065908 see the whole document	1-8, 10-14, 16-20, 22-26, 28-32

X Further documents are listed in the continuation of box C.	X Patent farrily members are listed in annex.
*Special categories of cited documents: *A* document defining the general state of the art which is not considered to be of particular relevance. *E* earlier document but published on or after the international filing date. *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified). *O* document referring to an oral disclosure, use, exhibition or other means. *P* document published prior to the international filing date but later than the priority date claimed.	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Oate of the actual completion of the international search 4 June 1998	Date of mailing of the International search report 2 4. 06, 98
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Klaver, T

Interna. .sal Application No PCT/US 98/00534

ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Κ	"candesartan cilexetil" DRUGS OF THE FUTURE, vol. 21, no. 7, 1996,	1-3, 10-12, 16-18,
	pages 738-739, XP002065909	22-24, 28-30
	see the whole document	
X	"candesartan cilexetil" DRUGS OF THE FUTURE, vol. 20, no. 7, 1995, pages 740-742, XP002065910	1-3, 10-12, 16-18, 22-24,
	see the whole document	28-30
x	T. LEJEMTEL ET AL.: "Irbesartan-a new angiotensin II antagonist: acute hemodynamic effects in patients with heart failure." CIRCULATION, vol. 94, no. 8 suppl., 1996, pages I622-I623, XP002065911 see the whole document	1-3, 10-12, 16-18, 22-24, 28-30
X	I. CROZIER ET AL.: "Losartan in heart failure" CIRCULATION , vol. 91, no. 3, 1995, pages 691-697, XP002065912 see the whole document	1-8, 10-14, 16-20, 22-26, 28-32
X	A. FUJIMORI ET AL.: "Effect of chronic treatment with YM358, an angiotensin II receptor (AT1) antagonist, in experimental heart failure." JPN. J. PHARMACOL., vol. 67, no. suppl. 1, 1995, page 124P XP002065913 see the whole document	1-3, 10-12, 16-18, 22-24, 28-30
X	T. KUSAYAMA ET AL.: "Effect of YM358, an angiotensin II receptor (AT1) antagost, on anesthetized dogs in acute left ventricular failure." JPN. J. PHARMACOL., vol. 67, no. suppl.1, 1995, page 124P XP002065914 see the whole document	1-3, 10-12, 16-18, 22-24, 28-30
A	Y. INADA ET AL.: "nonpeptide angiotensin II- receptor antagonists." FOLIA HARMACOL. JPN., vol. 104, no. 3, 1994, pages 217-228, XP002065915	
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C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A .	P. BÜHLMAYER ET AL.: "Valsartan, a potent, orally acie angiotensin II antagonist developed from the structurally new amino acid series." BIOORG. MED. CHEM. LETT., vol. 4, no. 1, 1994, pages 29-34, XP002065916	
A	B. NOËL ET AL: "Clinical and hormonal effects of the new angiotensin II receptor antagonist LRB081" J. CARDIOVASC. PHARMACOL., vol. 28, no. 2, 1996, pages 252-258, XP002065917	
A	WO 96 40257 A (SEARLE & CO) 19 December 1996	r
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International application No. PCT/US 98/00534

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 4-6,13,19,25,31 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of Item 2 of Iirst sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

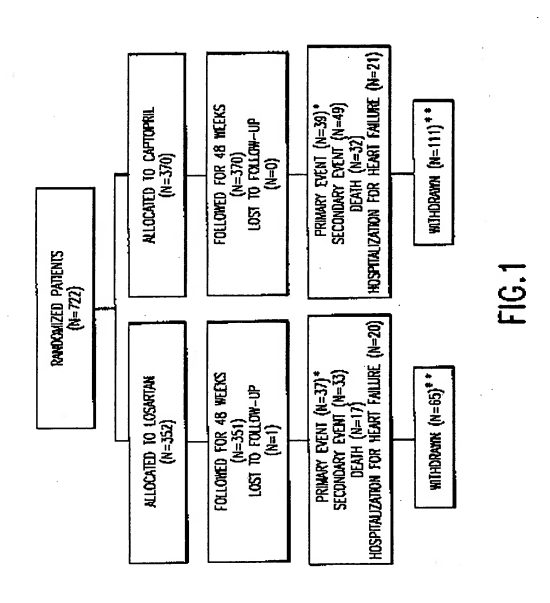
Claims Nos.: 4-6,13,19,25,31

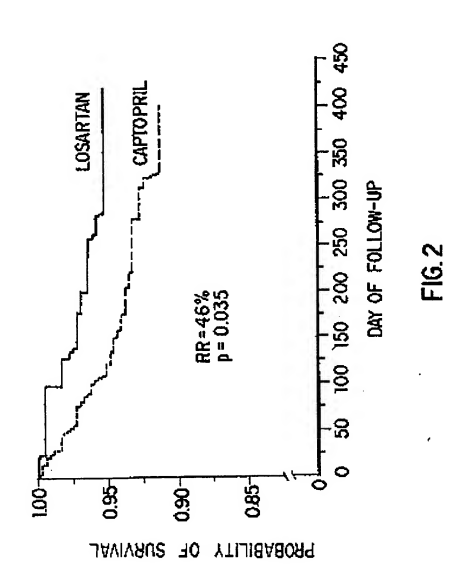
In view of the large number of compounds, which are defined by the general definition in the independent claims, the search had to be restricted for economic reasons. The search was limited to the compounds for which pharmacological data was given and/or the compounds mentioned in the claims, and to the general idea underlying the application. A considerable number of the acronyms used to identify the compounds could not be traced. As a consequence only the acronyms were used in the search, not the official names and synonyms. (see Guidelines, Chapter III, paragraph 2.3).

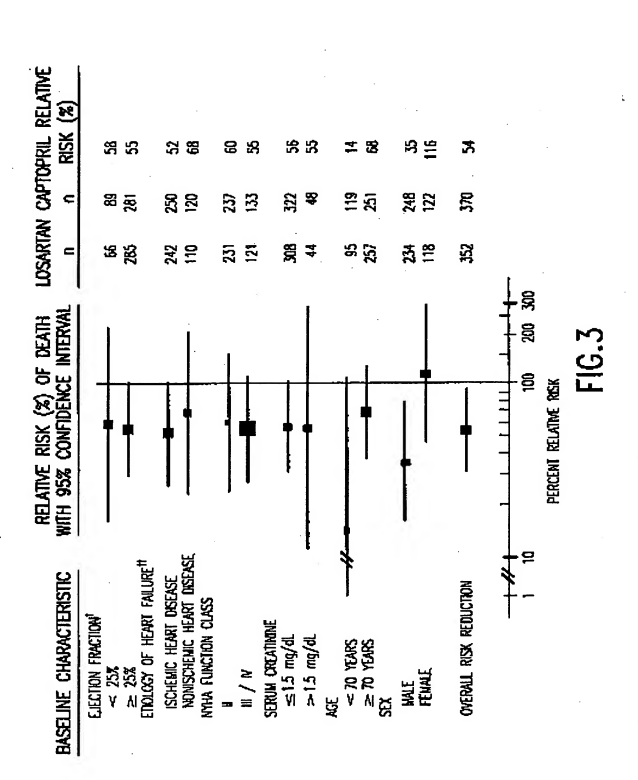
information on patent family members

Internation No PCT/US 98/00534

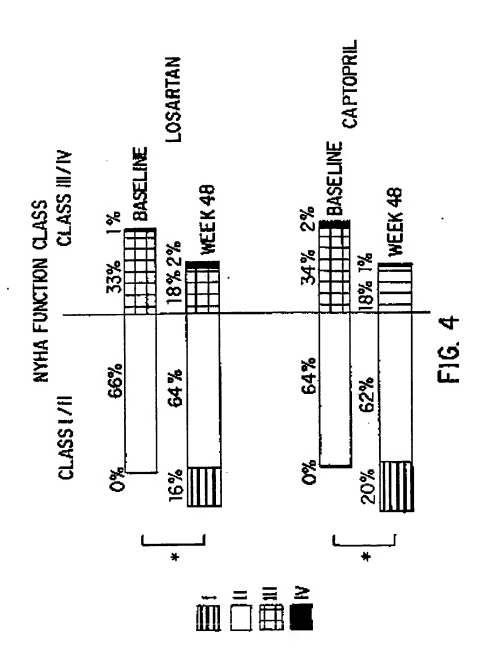
Patent document cited in search report		Publication date		atent family member(s)	Publication date
WO 9640257	A	19-12-1996	AU EP NO	6157796 A 0831910 A 975741 A	30-12-1996 01-04-1998 29-01-1998



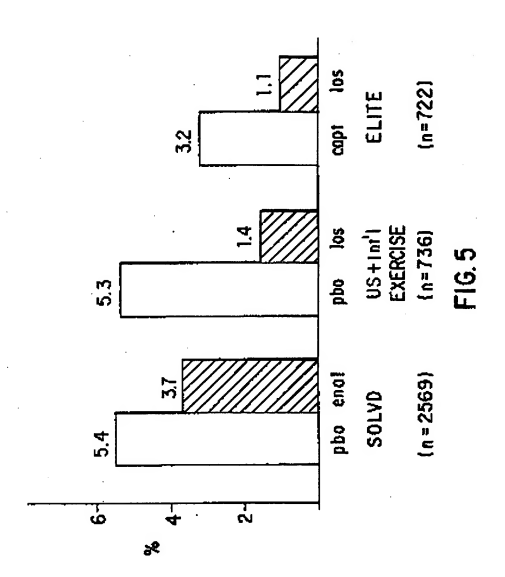




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